Respironics V60/V60 Plus Ventilator
User Manual
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Chapter 1. Warnings, cautions, and notes

Before using the Respironics V60/V60 Plus Ventilator on a patient, familiarize yourself with this user manual, particularly the safety considerations listed. Be aware, however, that this manual is a reference only. It is not intended to supersede your institution’s protocol regarding the safe use of assisted ventilation.

Definitions

| WARNING: | Alerts the user to the possibility of injury, death, or other serious adverse reactions associated with the use or misuse of the device. |
| CAUTION: | Alerts the user to the possibility of a problem with the device associated with its use or misuse, such as device malfunction, device failure, damage to the device, or damage to other property. |
| NOTE: | Emphasizes information of particular importance. |

General

| WARNING: | Always have immediate access to an alternative means of ventilation. If there is a ventilator failure, immediately remove the ventilator from use and secure an alternative means of ventilation, such as a self-inflating, manually powered resuscitator and mask. Failure to do so can result in patient injury or death. |
| WARNING: | Use the Respironics V60/V60 Plus Ventilator on spontaneously breathing patients only. It is an assist ventilator and is intended to augment the ventilation of a spontaneously breathing patient. It is not intended to provide the total ventilatory requirements of the patient. |
| WARNING: | We do not recommend you use the Respironics V60/V60 Plus Ventilator on patients who require ventilation at predetermined tidal volumes. The ventilator provides continuous positive airway pressure (CPAP) and positive pressure ventilation (S/T, PCV, AVAPS, and PPV) and is indicated for assisted ventilation only. These modes do not provide ventilation with guaranteed tidal volume delivery. |
Warnings, cautions, and notes

WARNING: We do not recommend you use AVAPS on patients who require rapid and frequent IPAP adjustments to maintain a consistent tidal volume. AVAPS, a volume targeted mode, changes the IPAP setting in order to achieve the target tidal volume. During AVAPS setup, there may be a period of time before the target tidal volume is achieved. AVAPS is ideal for more stabilized patients.

WARNING: To reduce the risk of CO₂ rebreathing, make sure EPAP pressures and exhalation times are sufficient to clear all exhaled gas through the exhalation port. In noninvasive ventilation continuous air flow through the port flushes exhaled gases from the circuit. The ability to completely exhaust exhaled gas from the circuit depends on the EPAP setting and I:E ratio. Higher tidal volumes further increase the volume of CO₂ rebreathed by the patient. Note: this may occur if exhalation time is insufficient.

WARNING: To reduce the risk of CO₂ rebreathing, monitor the patient for changes in respiratory status at the start of ventilation and with each change in ventilator settings, circuit configuration, or patient condition. Pay attention to ventilator alarms that warn of increased CO₂ rebreathing risk.

WARNING: To ensure accuracy of oxygen administration and to monitor for the presence of contamination (incorrect gas connected), use an external oxygen monitor to verify the oxygen concentration in the delivered gas.

WARNING: To reduce the risk of fire, use the ventilator in well-ventilated areas away from flammable anesthetics. Do not use in a hyperbaric chamber or other similarly oxygen-enriched environments. Do not use near an open flame.

WARNING: To reduce the risk of electric shock from liquid entering the device, do not put a container filled with a liquid on the ventilator.

WARNING: To reduce patient risk of oxygen toxicity, keep free-flowing oxygen away from air inlet of ventilator.

WARNING: The nurse call/remote alarm should be considered a backup to the ventilator's primary alarm system.

WARNING: Set the alarm loudness above the ambient level. Setting the alarm loudness too low may prevent recognition of alarm conditions.

WARNING: Avoid blocking the alarm speakers beneath the ventilator.

WARNING: Do not leave the ventilator unattended when stationed on an incline.

WARNING: The V60/V60 Plus Ventilator may cause radio interference or may disrupt the operation of nearby equipment. It may be necessary to take mitigation measures, such as re-orienting or relocating the ventilator or shielding the location.

WARNING: This Equipment is designed to comply with IEC 60601-1-2. This equipment generates, uses, and can radiate radio frequency energy and, if not installed and used in accordance with instructions, may cause harmful interference to other devices in the vicinity. However, there is no guarantee that interference will not occur in a particular installation. Harmful interference to other devices can be determined by turning this equipment ON and OFF. Try to correct the interference using one or more of the following:

- Reorient or relocate the receiving device
- Increase the separation between the equipment
Warnings, cautions, and notes

- Connect the equipment to an outlet on a different circuit from that to which the other device(s) are connected and consult the Philips service technician for help.
- Consult Philips for help.

WARNING: Use of non-approved accessories, transducers or cables may increase EMC emissions or decrease the EMC immunity performance of the equipment.

WARNING: Do not use the ventilator in an MRI environment. The V60/V60 Plus Ventilator is MR Unsafe. Keep it outside the MRI scan room (Zone IV). It represents a projectile hazard.

CAUTION: Federal law (USA) restricts this device to sale by or on the order of a physician.

CAUTION: The Respironics V60/V60 Plus Ventilator is designed to operate in the temperature range of 5 to 40 ºC (41 to 104 ºF). To minimize the risk of overheating the device, do not operate adjacent to heaters or other heat sources.

NOTE: The displays shown in this manual may not exactly match what you see on your own ventilator.

NOTE: Pressures are indicated on the ventilator in cmH2O. Millibars and hectopascals (hPa) are used by some institutions instead. Since 1 millibar equals 1 hPa, which equals 1.016 cmH2O, the units may be used interchangeably.

NOTE: The ventilator is not intended for use as an ambulance transport ventilator or as an Automatic Transport Ventilator as described by the American Hospital Association and referenced by the FDA. It is intended to allow the patient to be transported within the hospital setting using a cart to move the ventilator.

NOTE: When attachments or other components or subassemblies are added to the ventilator breathing system, the pressure gradient across the ventilator breathing system, measured with respect to the ventilator outlet, may increase.

NOTE: To ensure the correct performance of the ventilator and the accuracy of patient data, use only Respironics-approved accessories with the ventilator. See Appendix C, “Parts and accessories”.

NOTE: This Respironics V60/V60 Plus Ventilator and its recommended accessories that have patient contact are not made with natural rubber latex.

NOTE: If an alarm persists for no apparent reason, discontinue ventilator use and contact Philips.

NOTE: If you detect any unexplained changes in the performance or visual displays of the ventilator, discontinue ventilator use and contact Philips.

NOTE: The Respironics V60/V60 Plus Ventilator does not support automatic record keeping.
Warnings, cautions, and notes

NOTE: All ventilator mode and alarm settings, alarm messages and significant events are retained and automatically logged, even when power is lost.

NOTE: Any serious incident that has occurred in relation to this device should be reported to Philips and the competent authority of the country in which the user and/or patient is established.

Preparing for ventilation

WARNING: Connect the ventilator only to an appropriate medical-grade oxygen source.

WARNING: To reduce the risk of hypoxia, connect only oxygen to the high-pressure connector at the rear of the ventilator.

WARNING: The ventilator is a high-flow device. Connect it only to a gas supply system that can provide adequate flow to all terminal outlets. Connecting the ventilator to an adequate gas supply system helps ensure that the ventilator and other connected devices perform to their specifications.

WARNING: To reduce the risk of fire, do not use a high-pressure oxygen hose that is worn or contaminated with combustible materials like grease or oil.

WARNING: The Respironics V60/V60 Plus Ventilator is designed to use ambient air and high pressure 100% oxygen. No other gases should be used.

WARNING: Do not use the ventilator with helium or helium mixtures. The ventilator is not intended to be used with helium or heliox. Connecting helium to the ventilator may affect ventilator performance, gas mixtures, and measurements due to the lower density of helium.

WARNING: Do not use the ventilator with nitric oxide.

WARNING: To prevent possible asphyxia and to reduce the risk of CO₂ rebreathing, take these precautions with respect to mask and exhalation port use:

- Use only an oro-nasal mask with an anti-asphyxia valve or a nasal mask for noninvasive ventilation.
- Do not occlude the exhalation port.
- Turn on the ventilator and verify that the port is operational before application. Pressurized gas from the ventilator should cause a continuous flow of air to exhaust from the leak port, flushing exhaled gas from the circuit.
- Never leave the mask on the patient while the ventilator is not operating. When the ventilator is not operating, the exhalation port does not allow sufficient exhaust to eliminate CO₂ from the circuit. Substantial CO₂ rebreathing may occur.

WARNING: The patient’s exhaled volume can differ from the measured exhaled volume due to leaks around the mask during noninvasive ventilation.

WARNING: To ensure normal air circulation and exchange, do not cover or block the ports on the ventilator. Do not block the air inlet panel on the right side of the ventilator.
Warnings, cautions, and notes

WARNING: Do not cover or position the ventilator so as to adversely affect its operation or performance. For example, positioning the ventilator next to a curtain that blocks the flow of cooling air can cause the equipment to overheat. Use the V60/V60 Plus in an upright position that does not block the air inlet.

WARNING: Do not block the blower intake. Blocking the intake can impair ventilator performance and result in patient injury.

WARNING: To reduce the risk of the device overheating and possible burn injury, do not block the fan intake at the rear of the ventilator.

WARNING: To prevent possible patient injury and possible water damage to the ventilator, make sure the humidifier is set appropriately.

WARNING: When using a humidifier, always use either a circuit with a water trap or a heated wire circuit to minimize patient risk from condensate in the circuit.

WARNING: To prevent the possibility of inadequate humidification, pay close attention to the humidifier’s functioning when operating the ventilator at an ambient temperature > 30 ºC (86 ºF). The ventilator warms the air delivered to the patient above ambient temperature, which may impair the humidifier’s performance.

WARNING: To reduce the risk that the patient will aspirate condensed water from the breathing circuit, position any humidifier lower than both the ventilator and the patient.

WARNING: To prevent possible patient injury and equipment damage, do not turn the humidifier on until the gas flow has started and is regulated. Starting the heater or leaving it on without gas flow for prolonged periods may result in heat build-up, causing a bolus of hot air to be delivered to the patient. Circuit tubing may melt under these conditions. Turn the heater power switch off before stopping gas flow.

WARNING: To reduce the risk of fire, use only patient circuits intended for use in oxygen-enriched environments. Do not use antistatic or electrically conductive tubing.

WARNING: To prevent patient or ventilator contamination, always use a main flow bacteria filter on the patient gas outlet port. Filters not approved by Respironics may degrade system performance.

WARNING: During ventilation, patient exhale is released into room air. Use of a patient circuit with a filter on its exhalation port is recommended.

WARNING: To reduce the risk of bacterial contamination or damage, handle bacteria filters with care.

WARNING: Avoid adding any components to the patient circuit that are not absolutely necessary. Additional components installed in the patient circuit can change the pressure gradient across the ventilator breathing system, increase the dead space, and adversely affect the ventilator performance.

WARNING: Any additional accessories in the patient circuit may substantially increase flow resistance and impair ventilation.

WARNING: Avoid adding resistive circuit components on the patient side of the proximal pressure line. Such components may defeat the disconnect alarm.
Warnings, cautions, and notes

**WARNING:** To reduce the risk of strangulation from patient tubing, use a tubing support arm and secure the proximal pressure line with clips.

**WARNING:** To reduce the risk of electric shock, connect the ventilator to an AC supply mains with protective earth only.

**WARNING:** Do not use extension cords, adapters, or power cords with the ventilator that are not approved by Respironics.

**WARNING:** To prevent unintentional disconnection of the power cord, always use the correct, Philips-supplied power cord and lock it into place with the power cord retainer before you switch the ventilator on. The retainer is designed to hold the connector end of the Philips-supplied cord securely in place.

**WARNING:** The V60/V60 Plus Ventilator should not be positioned in a way that makes it difficult to disconnect from mains power if necessary. Disconnect from supply mains by removing the power cord from the wall outlet. The AC mains plug is used as disconnection device.

**WARNING:** To reduce the risk of electric shock, regularly inspect the AC power cord and verify that it is not frayed or cracked.

**WARNING:** To reduce the risk of strangulation, route the power cord to avoid entanglement.

**WARNING:** To reduce the risk of power failure to the ventilator, pay close attention to the battery’s charge level. The battery’s operation time is approximate and is affected by ventilator settings, discharge and recharge cycles, battery age, and ambient temperature.

**WARNING:** Always check the status of the oxygen cylinders before using the ventilator during transport.

**WARNING:** Provide external oxygen monitoring to minimize patient risk in case of O₂ supply loss or ventilator failure.

**WARNING:** To ensure the ventilator’s safe operation, always verify ventilator operation as described in “Verify ventilator operation” on page 5-8 before using the ventilator on a patient. If the ventilator fails any tests, remove it from clinical use immediately. Do not use the ventilator until necessary repairs are completed and all tests have passed.

**WARNING:** To prevent possible patient injury due to nonannunciating alarms, verify the operation of any remote alarm device before use.

**WARNING:** To prevent possible patient injury, always return alarm settings to hospital-standard values after verifying ventilator operation.

**WARNING:** Manufacturer default settings are not appropriate for all patients. Prior to using the ventilator, verify that the current alarm settings or defaults are appropriate for each particular patient.

**CAUTION:** To prevent possible damage to the ventilator, ensure that the connection to the oxygen supply is clean and unlubricated, and that there is no water in the oxygen supply gas.

**CAUTION:** For 120 V equipment, grounding reliability can only be achieved when it is connected to an equivalent receptacle marked “hospital only” or “hospital grade.”
Warnings, cautions, and notes

CAUTION: Oxygen hose configurations using SIS connectors generate higher resistance to flow. Therefore, a supply pressure of 53 to 87 psig is recommended when adding supplemental O2 accessories with SIS adapters such as the O2 transport manifold.

NOTE: The V60/V60 Plus Ventilator is a single-limb device with substantial intentional and unintentional leak in the ventilator breathing system. Under those conditions, CO2 cannot be measured accurately. Therefore, we do not recommend the use of CO2 monitoring.

Operation

WARNING: To prevent possible patient injury, avoid setting alarm limits to extreme values, which can render the alarm system useless.

WARNING: PPV limits are not intended to be the primary ventilator alarms and should not be substituted for the alarms found in the Alarm Settings window.

WARNING: To prevent the delivery of excessive pressure or volume, set the PPV limits appropriately. Delivery of excessive pressure or volume can occur from a sudden increase in mask leak, inappropriate settings, or a plugged or kinked proximal pressure line. Conversely, insufficient treatment may result if limits are set too low.

WARNING: Nebulization or humidification can increase the resistance of breathing system filters. When using a nebulizer or humidifier, monitor the breathing system filter frequently for increased resistance and blockage.

WARNING: Using a jet nebulizer can cause inadvertent alarms and affect the accuracy of delivered FiO2. To reduce patient risk, use only an approved nebulizer.

Operation in high flow therapy (HFT)

WARNING: When transitioning from a high flow therapy interface to an NIV mask, ensure that an exhalation port is placed in the circuit and is unobstructed to reduce the risk of CO2 rebreathing.

WARNING: When transitioning from ventilation to high flow therapy, remove the NIV mask and use only a Philips-approved high flow patient interface to minimize pressure build-up and patient discomfort.

WARNING: When transitioning from high flow therapy to ventilation, remove the high flow nasal cannula as it is restrictive and may defeat alarms such as patient disconnect. Using a high flow nasal cannula in an NIV mode may lead to hypercarbia due to the inability to provide pressure support.

WARNING: Patient alarms are not available during high flow therapy (HFT) as the therapy uses an open system. A high flow nasal cannula occupies only a portion of the nares and patients can breathe through their mouth, which prevents estimation of patient parameters such as tidal volume, respiratory rate, pressure, and minute ventilation. Provide external monitoring, including oximetry, to inform the clinician of a change in the patient’s condition.
Warnings, cautions, and notes

**WARNING:** During high flow therapy (HFT), verify that an occlusive patient interface is not being used. Occlusive patient interfaces include a cannula fully sealed within the nares, an NIV mask, or a direct connection to a tracheostomy tube or endotracheal tube. Remove any occlusive interface immediately as this may expose the patient to unintended high pressures.

Alarms and messages

**WARNING:** If AC power fails and the backup battery is not installed or is depleted, an audible and visual alarm annunciates for at least 2 minutes. Immediately discontinue ventilator use and secure an alternative means of ventilation. As in most ventilators with passive exhalation ports, when power is lost, sufficient air is not provided through the circuit and exhaled air may be rebreathed.

Care and maintenance

**WARNING:** To reduce the risk of electric shock, power down the ventilator and disconnect it from AC power before cleaning, disinfecting, or servicing it.

**WARNING:** Turn off the ventilator and disconnect it from the AC mains outlet before you perform decontamination or maintenance procedures. Failure to do so may result in electric shock.

**WARNING:** To prevent patient or ventilator contamination, inspect and replace the main flow bacteria filter between patients and at regular intervals (or as stated by the manufacturer).

**WARNING:** To prevent possible patient injury, inspect and verify the proper operation of the exhalation port regularly during use.

**WARNING:** To reduce the risk of fire, explosion, leakage, or other hazard, take these precautions with respect to the battery:

- Do not attempt to disassemble, open, drop, crush, bend or deform, insert foreign objects into, puncture, or shred the battery pack; modify or remanufacture it; immerse or expose it to water or other liquids; expose it to fire, excessive heat (including soldering irons); or put it in a microwave oven.

- Replace the battery only with another battery specified by the manufacturer.

- Follow all instructions for proper use of the battery.

- Do not short-circuit the battery or allow metallic or conductive objects to contact the battery connector housing.

- Use the battery with the Respironics V60/V60 Plus Ventilator only.

**WARNING:** Modification of the V60/V60 Plus Ventilator and associated equipment is not permitted and may compromise ventilator operation and patient safety. Service should only be performed by qualified service personnel.

**WARNING:** Only authorized service personnel should replace parts within the ventilator or perform other service activities. Unauthorized personnel without proper training are at risk of electric shock.
Warnings, cautions, and notes

**WARNING:** This product consists of devices that may contain mercury, which must be recycled or disposed of in accordance with local, state, or federal laws. (Within this system, the backlight lamps in the monitor display contain mercury.)

**CAUTION:** Do not attempt to sterilize or autoclave the ventilator.

**CAUTION:** To prevent possible damage to the ventilator, use only those cleaning and disinfecting agents listed in this manual.

**CAUTION:** To prevent possible damage to the ventilator, do not drip or spray any liquids directly onto any surface including the front panel, touchscreen, navigation ring (legacy versions), and Accept button.

**CAUTION:** Never clean or disinfect the touchscreen with an abrasive brush or device, since this will cause irreparable damage.

**CAUTION:** To avoid introducing foreign matter into the ventilator and to ensure proper system performance, change the air inlet filter at regular intervals (or as stipulated by your institution).

**CAUTION:** To ensure proper system performance, use a Respironics-approved air inlet filter.

**CAUTION:** Because some environments cause a quicker collection of lint and dust than others, inspect the filters more often when needed. The air inlet filter should be replaced; the cooling fan filter should be cleaned.

**CAUTION:** To prevent possible damage to the ventilator, always ship it with the original packing material. If the original material is not available, contact Philips to order replacements.

---

**First-time installation**

**WARNING:** Never attempt to disconnect or connect the battery during operation.

**CAUTION:** To prevent possible damage to the ventilator, always secure it to its stand or securely place it on a flat, stable surface that is free of dirt and debris. Do not use the ventilator adjacent to, or stack it with, other equipment.

---

**Communications interface**

**WARNING:** Connect to the ventilator only items that are specified as part of or compatible with the ventilator system. Additional equipment connected to medical electrical equipment must comply with the respective IEC or ISO standards. Furthermore, all configurations shall comply with the requirements for medical electrical systems (see IEC 60601-1-1 or clause 16 of edition 3 of IEC 60601-1, respectively). Anybody connecting additional equipment to medical electrical equipment configures a medical system and is therefore responsible for ensuring that the system complies with the requirements for medical electrical systems. Also be aware that local laws may take priority over the above mentioned requirements. If in doubt, consult Philips.

**WARNING:** The USB port is not currently available for use. DO NOT connect or attempt to power any equipment from the USB port.
Warnings, cautions, and notes

**WARNING:** It is the responsibility of the end user to validate the compatibility and use of information transmitted from the ventilator to the device to be connected to the ventilator.

**WARNING:** The data provided through the communications interface is for reference only. Decisions for patient care should be based on the clinician’s observations of the patient.

**WARNING:** To prevent possible patient injury due to nonannunciating alarms, verify the operation of any remote alarm device before use.

**WARNING:** To ensure the functionality of the remote alarm, connect only Respironics-approved cables to the remote alarm port.

**CAUTION:** The remote alarm port is intended to connect only to an SELV (safety extra-low voltage and ungrounded system with basic insulation to ground), in accordance with IEC 60601-1. To prevent damage to the remote alarm, make sure the signal input does not exceed the maximum rating of 24 VAC or 36 VDC at 500 mA with a minimum current of 1 mA.

---

**Diagnostic mode**

**WARNING:** To prevent possible patient injury, do not enter the diagnostic mode while a patient is connected to the ventilator. Verify that the patient is disconnected before proceeding.
Chapter 2. Symbols

Refer to these tables to interpret symbols used on the ventilator labels, backup battery labels, and packaging and on the ventilator screen. To interpret symbols pertaining to accessories, refer to their instructions for use.

Table 2-1: Symbols used on ventilator labels, battery, and packaging

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="Warning.png" alt="Warning" /></td>
<td>Warning: Risk of explosion. Do not use in the presence of flammable anesthetics.</td>
</tr>
<tr>
<td><img src="Attention.png" alt="Attention" /></td>
<td>Attention, consult the accompanying documents.</td>
</tr>
<tr>
<td><img src="Read.png" alt="Read" /></td>
<td>Read the user manual before using the ventilator. Symbol may be accompanied by the web address <a href="http://www.Philips.com/IFU">www.Philips.com/IFU</a> to indicate access to electronic IFUs.</td>
</tr>
<tr>
<td><img src="Electronic.png" alt="Electronic" /></td>
<td>Electronic instructions for use. Indicates that relevant information for use of the product is available in electronic form.</td>
</tr>
<tr>
<td><img src="Mandatory.png" alt="Mandatory" /></td>
<td>(Blue) It is mandatory for the operator to consult the accompanying documents.</td>
</tr>
<tr>
<td><img src="Protective.png" alt="Protective" /></td>
<td>Protective earth (ground)</td>
</tr>
<tr>
<td><img src="Medical.png" alt="Medical" /></td>
<td>Medical device</td>
</tr>
<tr>
<td><img src="UDI.png" alt="UDI" /></td>
<td>Unique device identifier</td>
</tr>
<tr>
<td><img src="Distributor.png" alt="Distributor" /></td>
<td>Distributor. Symbol accompanied by address</td>
</tr>
<tr>
<td><img src="Importer.png" alt="Importer" /></td>
<td>Importer. Symbol accompanied by address</td>
</tr>
</tbody>
</table>
Symbols

Table 2-1: Symbols used on ventilator labels, battery, and packaging

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Fragile" /></td>
<td>Fragile</td>
</tr>
<tr>
<td><img src="image" alt="Keep dry" /></td>
<td>Keep dry</td>
</tr>
<tr>
<td><img src="image" alt="Stacking limit by number" /></td>
<td>Stacking limit by number</td>
</tr>
<tr>
<td><img src="image" alt="This end up" /></td>
<td>This end up</td>
</tr>
<tr>
<td><img src="image" alt="Type B applied part" /></td>
<td>Type B applied part, which is equipment that provides a particular degree of protection against electric shock, particularly in regard to allowable leakage current and of the protective earth connection</td>
</tr>
<tr>
<td><img src="image" alt="Requires alternating current (AC)" /></td>
<td>Requires alternating current (AC)</td>
</tr>
<tr>
<td><img src="image" alt="IPX1" /></td>
<td>Degree of fluid ingress protection provided by the enclosure (drip-proof)</td>
</tr>
<tr>
<td><img src="image" alt="IP21" /></td>
<td>Degree of solid object protection and fluid ingress protection provided by the enclosure (drip-proof)</td>
</tr>
<tr>
<td><img src="image" alt="Rx ONLY" /></td>
<td>Caution: Federal law restricts this device to sale by or on the order of a physician</td>
</tr>
<tr>
<td><img src="image" alt="Alarm and remote alarm" /></td>
<td>Alarm and remote alarm</td>
</tr>
<tr>
<td><img src="image" alt="Two states of control: ON and Shutdown" /></td>
<td>Two states of control: ON and Shutdown</td>
</tr>
<tr>
<td><img src="image" alt="Battery" /></td>
<td>Battery</td>
</tr>
<tr>
<td><img src="image" alt="European Conformity. Symbol is on rear panel of ventilator." /></td>
<td>European Conformity. Symbol is on rear panel of ventilator.</td>
</tr>
</tbody>
</table>
Table 2-1: Symbols used on ventilator labels, battery, and packaging

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Brazilian Conformity" /></td>
<td>Brazilian Conformity. Certification by INMETRO (National Institute of Metrology, Standardization and Industrial Quality)/SGS (Societe Generale de Surveillance). One of these three symbols, depending upon available space.</td>
</tr>
<tr>
<td><img src="image" alt="EAC" /></td>
<td>EurAsian Conformity mark - EAC</td>
</tr>
<tr>
<td><img src="image" alt="Date of manufacture" /></td>
<td>Date of manufacture. Symbol accompanied by date.</td>
</tr>
<tr>
<td><img src="image" alt="Country of Manufacture" /></td>
<td>Country of Manufacture. Symbol accompanied by country code (and optionally manufacture date and/or manufacturer’s name and address).</td>
</tr>
<tr>
<td><img src="image" alt="Manufacturer" /></td>
<td>Manufacturer. Symbol accompanied by manufacturer’s name and address.</td>
</tr>
<tr>
<td><img src="image" alt="EC representative" /></td>
<td>EC representative</td>
</tr>
<tr>
<td><img src="image" alt="Serial number" /></td>
<td>Serial number</td>
</tr>
<tr>
<td><img src="image" alt="Order number" /></td>
<td>Order number</td>
</tr>
<tr>
<td><img src="image" alt="Lot or batch number" /></td>
<td>Lot or batch number</td>
</tr>
<tr>
<td><img src="image" alt="Model number" /></td>
<td>Model number</td>
</tr>
<tr>
<td><img src="image" alt="Use by date" /></td>
<td>Use by date</td>
</tr>
<tr>
<td><img src="image" alt="RS-232 serial input/output" /></td>
<td>RS-232 serial input/output</td>
</tr>
<tr>
<td><img src="image" alt="USB port" /></td>
<td>USB port</td>
</tr>
</tbody>
</table>
Table 2-1: Symbols used on ventilator labels, battery, and packaging

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$O_2$</td>
<td>Oxygen</td>
</tr>
<tr>
<td>⚠️</td>
<td>(Yellow) Warning</td>
</tr>
<tr>
<td>Ethernet connection</td>
<td></td>
</tr>
<tr>
<td>🔄</td>
<td>Accept button on the top-right front of the ventilator</td>
</tr>
<tr>
<td>⏬</td>
<td>Adjustment direction on the navigation ring (legacy versions only)</td>
</tr>
<tr>
<td>🇨🇦</td>
<td>Canadian Standards Association approval</td>
</tr>
<tr>
<td>✘</td>
<td>Do not disassemble. Refer to authorized service personnel.</td>
</tr>
<tr>
<td>🚫</td>
<td>Product must be disposed of in accordance with the WEEE directive.</td>
</tr>
<tr>
<td>🎭</td>
<td>Noninvasive ventilation (patient with mask)</td>
</tr>
<tr>
<td>🎭</td>
<td>Invasive ventilation (intubated patient)</td>
</tr>
<tr>
<td>✘</td>
<td>The V60/V60 Plus Ventilator is MR Unsafe and presents a projectile hazard. Keep the ventilator outside MRI scan room (Zone IV).</td>
</tr>
</tbody>
</table>
### Symbols

Table 2-1: Symbols used on ventilator labels, battery, and packaging

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image1.png" alt="Symbol" /></td>
<td>Do not block the cooling fan Inlet (at the rear of the ventilator).</td>
</tr>
<tr>
<td><img src="image2.png" alt="Symbol" /></td>
<td>Used in conjunction with the blue “consult accompanying documents” symbol to indicate “Do not block the cooling fan Inlet.”</td>
</tr>
<tr>
<td><img src="image3.png" alt="Symbol" /></td>
<td>No pushing. Do not push on the ventilator screen. Tipping hazard.</td>
</tr>
<tr>
<td><img src="image4.png" alt="Symbol" /></td>
<td>Total mass (weight). See page 11-6 for details on physical characteristics of the ventilator, stand, and accessories.</td>
</tr>
<tr>
<td><img src="image5.png" alt="Symbol" /></td>
<td>Hospital-grade (On power cord)</td>
</tr>
<tr>
<td><img src="image6.png" alt="Symbol" /></td>
<td>Packaging unit. Symbol accompanied by a number (indicating the number of pieces in the package)</td>
</tr>
<tr>
<td><img src="image7.png" alt="Symbol" /></td>
<td>Recycle</td>
</tr>
<tr>
<td><img src="image8.png" alt="Symbol" /></td>
<td>Recycle (Taiwan)</td>
</tr>
<tr>
<td><img src="image9.png" alt="Symbol" /></td>
<td>RoHS (China). Administrative Measure on the Control of Pollution Caused by Electronic Information Products. Contains RoHS substances with 50 years environmentally friendly use period (EFUP).</td>
</tr>
<tr>
<td><img src="image10.png" alt="Symbol" /></td>
<td>uR UL recognition symbol</td>
</tr>
<tr>
<td><img src="image11.png" alt="Symbol" /></td>
<td>Direct current (DC). Symbol is on backup battery.</td>
</tr>
<tr>
<td><img src="image12.png" alt="Symbol" /></td>
<td>Battery check.</td>
</tr>
</tbody>
</table>
Symbols

Table 2-1: Symbols used on ventilator labels, battery, and packaging

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Symbol" /></td>
<td>Rechargeable battery. Symbol is on backup battery.</td>
</tr>
<tr>
<td><img src="image" alt="Symbol" /></td>
<td>Lithium-ion battery. Battery must be recycled or disposed of properly. Symbol is on backup battery.</td>
</tr>
<tr>
<td><img src="image" alt="Symbol" /></td>
<td>Barometric pressure limitation. Indicates the acceptable upper and lower limits of barometric pressure for transport and storage.</td>
</tr>
<tr>
<td><img src="image" alt="Symbol" /></td>
<td>Humidity limitation. Indicates the acceptable upper and lower limits of relative humidity for transport and storage.</td>
</tr>
<tr>
<td><img src="image" alt="Symbol" /></td>
<td>Temperature limit. Indicates the maximum and minimum temperature limits at which the item shall be stored or transported.</td>
</tr>
<tr>
<td><img src="image" alt="Symbol" /></td>
<td>Battery included</td>
</tr>
<tr>
<td><img src="image" alt="Symbol" /></td>
<td>C-Flex feature</td>
</tr>
<tr>
<td><img src="image" alt="Symbol" /></td>
<td>AVAPS mode (included)</td>
</tr>
<tr>
<td><img src="image" alt="Symbol" /></td>
<td>PPV software option</td>
</tr>
<tr>
<td><img src="image" alt="Symbol" /></td>
<td>Auto-Trak+ software option</td>
</tr>
<tr>
<td><img src="image" alt="Symbol" /></td>
<td>High flow therapy</td>
</tr>
</tbody>
</table>

**Note:** 3.00 software and above. HFT is optional for model V60 and included with model V60 Plus.
### Table 2-2: Symbols used on graphical user interface

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="" alt="Alarm (audible)" /></td>
<td>Alarm (audible)</td>
</tr>
<tr>
<td><img src="image" alt="Alarm is silenced" /></td>
<td>Alarm is silenced</td>
</tr>
<tr>
<td><img src="image" alt="High priority alarm" /></td>
<td>High priority alarm</td>
</tr>
<tr>
<td><img src="image" alt="Low priority alarm" /></td>
<td>Low priority alarm</td>
</tr>
<tr>
<td><img src="image" alt="Alarm reset" /></td>
<td>Alarm reset</td>
</tr>
<tr>
<td><img src="image" alt="Informational message" /></td>
<td>Informational message</td>
</tr>
<tr>
<td><img src="image" alt="Alarm message is displayed" /></td>
<td>Alarm message is displayed. Touch to hide alarm messages.</td>
</tr>
<tr>
<td><img src="image" alt="Alarm message is hidden" /></td>
<td>Alarm message is hidden. Touch to display alarm messages.</td>
</tr>
<tr>
<td><img src="image" alt="Do not use an NIV mask during high flow therapy" /></td>
<td>Do not use an NIV mask during high flow therapy (3.00 software and above, and V60 Plus).</td>
</tr>
<tr>
<td><img src="image" alt="Increase and decrease" /></td>
<td>Increase and decrease (adjustment arrow) buttons. Adjusts a setting or selects a value.</td>
</tr>
<tr>
<td><img src="image" alt="Accept button" /></td>
<td>Accept button. Accepts set values.</td>
</tr>
<tr>
<td><img src="image" alt="Cancel button" /></td>
<td>Cancel button. Cancels set values.</td>
</tr>
<tr>
<td><img src="image" alt="+2:00 minutes button" /></td>
<td>+2:00 minutes button. Adds two minutes to 100% $O_2$ delivery.</td>
</tr>
</tbody>
</table>
### Symbols

Table 2-2: Symbols used on graphical user interface (continued)

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image1" alt="Battery-AC" /></td>
<td>Ventilator is powered by AC power and the battery is installed.</td>
</tr>
<tr>
<td><img src="image2" alt="Battery-AC-IDLE" /></td>
<td>Ventilator is powered by AC power and the battery is not installed.</td>
</tr>
<tr>
<td><img src="image3" alt="Battery-Battery" /></td>
<td>Ventilator is powered by the battery. This symbol shows the approximate battery time remaining in hours and minutes, and it shows the capacity graphically.</td>
</tr>
<tr>
<td><img src="image4" alt="Help" /></td>
<td>Help button. Touch to display onscreen help information.</td>
</tr>
<tr>
<td><img src="image5" alt="Vertical Autoscale" /></td>
<td>Vertical autoscale button. Autoscales the Y axis of the graphs to fit the data currently displayed.</td>
</tr>
<tr>
<td><img src="image6" alt="Pause" /></td>
<td>Pause button. Freezes waveforms in the Waveform window.</td>
</tr>
<tr>
<td><img src="image7" alt="Pause in progress" /></td>
<td>Pause in progress</td>
</tr>
<tr>
<td><img src="image8" alt="Resume" /></td>
<td>Resume button. Resumes all waveform graphs from a paused state.</td>
</tr>
<tr>
<td><img src="image9" alt="Time Base Adjust" /></td>
<td>Time base adjust button. Rescales the X axis of the graph display data at 3, 6, 12, and 24 second increments.</td>
</tr>
</tbody>
</table>

- $V_E$: Estimated minute ventilation
- $V_T$: Estimated exhaled tidal volume
- $T_I/T_{TOT}$: Duty cycle. Inspiratory time divided by total cycle time.
### Symbols

Table 2-2: Symbols used on graphical user interface (continued)

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>![Symbol]</td>
<td>No valid data to display</td>
</tr>
<tr>
<td>![Symbol]</td>
<td>Data is under range</td>
</tr>
<tr>
<td>![Symbol]</td>
<td>Data is over range</td>
</tr>
<tr>
<td>![Symbol]</td>
<td>Pressure, centimeters of water</td>
</tr>
<tr>
<td>![Symbol]</td>
<td>Flow, liters per minute. BTPS compensated.</td>
</tr>
<tr>
<td>![Symbol]</td>
<td>Volume, milliliters</td>
</tr>
<tr>
<td>![Symbol]</td>
<td>Ramp Time is OFF (no ramp time set).</td>
</tr>
<tr>
<td>![Symbol]</td>
<td>Intentional leak. The number corresponds to the leak symbol printed on Philips Respironics masks.</td>
</tr>
</tbody>
</table>
Symbols
Chapter 3. General information

This manual covers the Respironics V60 and V60 Plus Ventilator configurations. Both share the same platform. The V60 Plus Ventilator comes standard with High Flow Therapy (HFT). The V60 Ventilator can be field-upgraded with HFT, subject to local regulations. For a full list of features, modes, and options, see "General description" on page 3-3.

NOTE: The 3.00 software upgrade, which permits the activation of HFT, and the V60 Plus Ventilator are not available in all countries.

Intended use

The Respironics V60/V60 Plus ventilator is an assist ventilator and is intended to augment patient breathing. It is intended for spontaneously breathing individuals who require mechanical ventilation: patients with respiratory failure, chronic respiratory insufficiency, or obstructive sleep apnea in a hospital or other institutional settings under the direction of a physician.

The Respironics V60/V60 Plus ventilator is intended to support pediatric patients (children and adolescents, weighing 20 Kg (44 lb) or greater), and adult patients. It is also intended for intubated patients meeting the same selection criteria as the noninvasive applications. The ventilator is intended to be used by qualified medical professionals, such as physicians, nurses, and respiratory therapists. The ventilator is intended to be used only with various combinations of Philips-recommended patient circuits, interfaces (masks), humidifiers, and other accessories.

Indications for use

The Respironics V60/V60 Plus is an assist ventilator and is indicated for use to augment patient breathing. The ventilator is indicated for spontaneously breathing individuals who require mechanical ventilation: patients with respiratory failure, chronic respiratory insufficiency, or obstructive sleep apnea in a hospital or other institutional settings under the direction of a physician.

Patient population

The Respironics V60/V60 Plus ventilator is intended to support pediatric patients (children and adolescents, weighing 20 kg (44 lb.) or greater), and adult patients. The ventilator is also intended for intubated patients meeting the same selection criteria as the noninvasive applications.
General information

Contraindications
The Respironics V60/V60 Plus Ventilator is contraindicated for patients with any of the following conditions:

- Lack of spontaneous respiratory drive
- Inability to maintain a patent airway or adequately clear secretions
- At risk for aspiration of gastric contents
- Acute sinusitis or otitis media
- Hypotension
- Untreated pertussis
- Epistaxis (nosebleed)

About CO₂ rebreathing
As with mask ventilation in general, patient CO₂ rebreathing may occur under some circumstances. Follow these guidelines to minimize the potential for CO₂ rebreathing. If rebreathing is a significant concern for a particular patient and these guidelines are not sufficient to acceptably reduce the potential for CO₂ rebreathing, consider an alternative means of ventilation.

- Increase EPAP to decrease the potential for CO₂ rebreathing. Higher pressures produce more flow through the exhalation port, which helps to purge all CO₂ from the circuit to prevent rebreathing.
- Be aware that the potential for CO₂ rebreathing increases as inspiratory time increases. A longer inspiratory time decreases exhalation time, allowing less CO₂ to be purged from the circuit before the next cycle. In such circumstances, higher tidal volumes further increase the volume of CO₂ rebreathed by the patient.

Potential side effects
Advise the patient to immediately report any unusual chest discomfort, shortness of breath, or severe headache. Other potential side effects of noninvasive positive pressure ventilation include: ear discomfort, conjunctivitis, skin abrasions due to mask/patient interface, and gastric distention (aerophagia). If skin irritation or breakdown develops from the use of the mask, refer to the accompanying mask instructions for appropriate action.
General description

The Respironics V60/V60 Plus Ventilator (Figure 3-1) is a microprocessor-controlled, bilevel positive airway pressure (BiPAP) ventilatory assist system that provides noninvasive positive pressure ventilation (NPPV) and invasive ventilatory support for spontaneously breathing adult and pediatric patients.

Ventilation modes. The ventilator offers a range of conventional pressure modes, CPAP (continuous positive airway pressure), PCV (pressure-controlled ventilation), and S/T (spontaneous/timed). The volume-targeted AVAPS (average volume-assured pressure support) mode combines the attributes of pressure-controlled and volume-targeted ventilation. The optional PPV mode provides pressure ventilation in proportion to the patient’s efforts.

Modes, therapies and features. Table 3-1 shows which modes, therapies and features are included or optional for the V60 and V60 Plus models.

Table 3-1: V60 and V60 Plus comparison

<table>
<thead>
<tr>
<th>Ventilator Model</th>
<th>Modes</th>
<th>Therapy</th>
<th>Features</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>AVAPS</td>
<td>PPV</td>
<td>HFT</td>
</tr>
<tr>
<td>V60</td>
<td>Included</td>
<td>Optional</td>
<td>Optional</td>
</tr>
<tr>
<td>V60 Plus</td>
<td>Included</td>
<td>Optional</td>
<td>Included</td>
</tr>
</tbody>
</table>

High flow therapy (HFT). High flow therapy provides a set flow of mixed air and oxygen. Flow and O2 percentage settings are selected by the clinician. HFT is available for 3.00 software and above, as well as for the V60 Plus.

Auto-Trak Sensitivity allows the ventilator to automatically compensate for intentional and unintentional leaks by maintaining a stable baseline and adjusting trigger and cycle thresholds for optimum patient-to-ventilator synchrony. The optional Auto-Trak+ feature lets you further adjust the level of Auto-Trak Sensitivity.

User interface. The ventilator’s 12.1-inch (31-cm) color touchscreen, Accept button, navigation ring (legacy versions only), and key panel let you easily access ventilator settings and monitored parameters.
General information

**Monitoring.** The ventilator displays monitored parameters as numbers and as real-time waveforms (curves or scalars).

**Alarms.** The ventilator’s operator-adjustable and nonadjustable alarms help ensure the patient’s safety.

**Power and gas supplies.** The ventilator uses AC mains as its primary power source. An internal backup battery provides a secondary power source. For more information, see Table 11-10 on page 11-8.

The ventilator uses high-pressure oxygen. An integral blower pressurizes gas for delivery to the patient.

---

**NOTE:** Oxygen delivered through the compressed gas hose and blower is used as fresh gas.

**Mounting.** The ventilator can be mounted to a stand. When equipped with the optional cylinder holder, the stand can accommodate two E-size oxygen cylinders. An oxygen manifold kit is available, which allows two oxygen cylinders and one wall oxygen supply line to be used as inputs to the ventilator.

**Communications interface.** The ventilator can output data through the RS-232 serial port upon receiving a command from a host computer or bedside monitoring system. The ventilator is equipped with a remote alarm/nurse call connection to activate alarms remotely.

**Upgradability via Respi-Link remote diagnostic system.** The Respi-Link interface permits software upgrade and remote troubleshooting of the ventilator through the RS-232 port.

Physical description

**Patient circuits, masks/patient interfaces, and accessories**

Figure 3-2 shows the Respironics V60/V60 Plus Ventilator with its patient circuit and accessories. Appendix C provides ordering information for parts and accessories.
General information

Figure 3-2: Respironics V60/V60 Plus Ventilator with accessories
General information

Ventilator unit

Depending on the version of the ventilator you have, the Accept button on the upper-right front of the device may or may not be surrounded by a navigation ring (scrollable wheel).

Newer versions have a streamlined appearance and the Accept button, and look like this:

Legacy versions have both the navigation ring and Accept button, and look like this:

The photos and illustrations throughout this manual reflect the newer version of the ventilator (Accept button without navigation ring). For more information on the functionality differences, see "Navigating the graphical user interface" on page 3-13.

Figure 3-3 through Figure 3-5 show the controls, indicators, and other important parts of the ventilator unit.
**General information**

![Figure 3-3: Front view](image)

<table>
<thead>
<tr>
<th>Number</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td><strong>Graphical user interface.</strong> Color LCD (liquid crystal display) with touchscreen.</td>
</tr>
<tr>
<td>2</td>
<td><strong>Accept button.</strong> Activates selections.</td>
</tr>
<tr>
<td>3</td>
<td><strong>Proximal pressure port.</strong> Connection for tubing that monitors patient pressure in the patient circuit.</td>
</tr>
<tr>
<td>4</td>
<td><strong>Ventilator outlet (To patient) port.</strong> Main connection for the patient circuit. Delivers air and oxygen in prescribed pressures to the patient.</td>
</tr>
<tr>
<td>5</td>
<td><strong>Alarm speakers</strong> (beneath ventilator)</td>
</tr>
<tr>
<td>6</td>
<td><strong>Alarm LED.</strong> Flashes during a high-priority alarm. On continuously during a ventilator inoperative condition.</td>
</tr>
<tr>
<td>7</td>
<td><strong>Battery (charged) LED.</strong> Flashes when battery is charging. On continuously when battery is charged. Off when ventilator is running on battery, when a battery error or failure is detected, or when the ventilator is off and AC power is not connected.</td>
</tr>
<tr>
<td>8</td>
<td><strong>ON/Shutdown key with LED.</strong> Turns on AC power and initiates ventilator shutdown. LED is continuously on when AC power is connected.</td>
</tr>
</tbody>
</table>
General information

Figure 3-4: Side view

<table>
<thead>
<tr>
<th>Number</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td><strong>Ventilation vents.</strong> Allow intake of air for delivery to the patient.</td>
</tr>
<tr>
<td>2</td>
<td><strong>Air inlet filter</strong> (under side panel). Filters the air for delivery to the patient.</td>
</tr>
</tbody>
</table>
### General information

![Figure 3-5: Rear view](image)

<table>
<thead>
<tr>
<th>Number</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Backup battery (compartment under side panel).</td>
</tr>
<tr>
<td>2</td>
<td>Remote alarm/nurse call connector</td>
</tr>
<tr>
<td>3</td>
<td>Reserved for future use</td>
</tr>
<tr>
<td>4</td>
<td>Power cord retainer</td>
</tr>
<tr>
<td>5</td>
<td>Power cord</td>
</tr>
<tr>
<td>6</td>
<td><strong>RS-232 serial connector (female DB-25).</strong> Connects to hospital information systems and other serial devices. Connects Respi-Link remote diagnostic system gateway for software updates.</td>
</tr>
<tr>
<td>7</td>
<td>Cooling fan filter</td>
</tr>
<tr>
<td>8</td>
<td>High-pressure oxygen inlet connector</td>
</tr>
<tr>
<td>9</td>
<td>Option labels</td>
</tr>
</tbody>
</table>
WARNING: To reduce the risk of power failure to the ventilator, pay close attention to the battery's charge level. The battery's operation time is approximate and is affected by ventilator settings, discharge and recharge cycles, battery age, and ambient temperature.

NOTE: The backup batteries are intended for short-term use only. They are not intended to be a primary power source.

NOTE: We recommend that the ventilator's batteries be fully charged before you ventilate a patient. If the batteries are not fully charged and AC power fails, always pay close attention to the level of battery charge.

NOTE: A new backup battery should be installed and charged within one year of the date of manufacture identified on the battery and on the shipping box.

NOTE: Storing a battery for an extended period of time after its manufacture date without being put into service, or at temperatures that exceed its limits, increases the risk of discharge whereby the battery is unable to be recharged by the ventilator.

NOTE: Batteries that are not installed in the ventilator should be stored at -20 to 25ºC and charged at least once per year.

NOTE: Lithium Ion batteries are shipped with a charge level of under 30%, which may further impact the time until the battery is charged and discharged.

The internal backup battery protects the ventilator from low, or failure of, AC (mains) power. If AC power fails, the ventilator automatically switches to operation on backup battery with no interruption in ventilation. The battery powers the ventilator until AC power is again adequate or until the battery is depleted. For electrical specifications, see Table 11-10 on page 11-8.

As a safeguard, the ventilator provides a low battery alarm. It also has a capacitor-driven backup alarm that sounds for at least 2 minutes when battery power is completely lost.

The ventilator charges the battery whenever the ventilator is connected to AC, with or without the ventilator switched on. The Battery (charged) LED flashes to show that the battery is being charged.

Check the battery charge level before putting a patient on the ventilator and before unplugging the ventilator for transport or other purposes. The power source symbol at the bottom right-hand corner of the screen shows the power source in use and, if the ventilator is running on battery, the level of battery charge (Figure 3-6).

NOTE: The battery charge level displays within about a minute of power on.

If the battery is not fully charged, recharge it by connecting the ventilator to AC power for a minimum of 5 hours. Pressing the Help button shows you the approximate time remaining until the battery is fully charged. If the battery is not fully charged within 16 hours, or the ventilator displays a Check Vent: Battery Failed alarm, replace the battery.
CAUTION: Avoid allowing the ventilator battery to become completely discharged. Otherwise, the battery may become over-discharged and require long recharge times of up to 16 hours or more. The over-discharged condition may permanently damage the battery so that it is unable to recharge. To prevent the occurrence of a non-recoverable over-discharged battery, always keep the ventilator connected to an AC outlet.

Figure 3-6: Power indicators
General information

About the graphical user interface

Through the graphical user interface (Figure 3-7) you make ventilator settings and view ventilator and patient data. During ventilation, the upper screen displays alarms and patient data. The middle screen displays real-time waveforms and alarm and informational messages. The lower screen lets you access modes and other ventilator settings, display help information, and see the power status.

Alarm status bar (see page 9-2)

Patient data window (see page 8-1)

Waveforms window (see page 8-1)

100% O₂ button (see page 6-6)

Window/window tabs (see page 6-1)

15% O2 button (see page 6-6)

Figure 3-7: Parts of graphical user interface
Navigating the graphical user interface

Select a function by touching the desired tab or button on the touchscreen.

If your ventilator has only the Accept button on the top-right front of the device (newer versions), you adjust values and navigate the graphical user interface by using the touchscreen.

If your ventilator has a navigation ring around the Accept button on the top-right front of the device (legacy versions), you can adjust values and navigate the graphical user interface by either rotating your finger on the navigation ring or by using the touchscreen.

<table>
<thead>
<tr>
<th>Touchscreen navigation</th>
<th>Front panel equivalent</th>
</tr>
</thead>
<tbody>
<tr>
<td>➤ Touch increase button (adjustment arrow). Press and hold for faster adjustments.*</td>
<td>On navigation ring (legacy versions), touch and rotate finger clockwise to increase value or move cursor forward</td>
</tr>
<tr>
<td>➥ Touch decrease button (adjustment arrow). Press and hold for faster adjustments.*</td>
<td>On navigation ring (legacy versions), touch and rotate finger counterclockwise to decrease value or move cursor backward</td>
</tr>
<tr>
<td>☑️ Touch Accept button to apply selection</td>
<td>Press Accept (checkmark) button to apply selection</td>
</tr>
</tbody>
</table>

* Available in Revision 2.30 software and above.

After making selections and adjusting values, accept those selections and apply the changes by either touching the Accept button in the user interface or by pressing the Accept button (checkmark) on the top-right front of the ventilator.

**To open a window**, touch the window tab.

**To cancel a function and close the window**, either select Cancel or touch another window tab.

**To adjust a parameter**, touch the arrow button or select the value with the navigation ring (legacy versions). Each touch changes the value in single increments or, for parameters with wide ranges, press and hold the arrow key to make faster changes. The slider flag moves along the setting range scale. Select Accept to apply.
The navigation ring (on legacy versions) also lets you adjust the position of the cursor in the waveforms window while the screen is frozen. See "Freezing and unfreezing waveforms" on page 8-3 for more information.
Starting up the ventilator

NOTE: Upon power-on the ventilator automatically runs a test of the backup audible alarm followed by the primary audible alarm. You should hear a high-pitched tone, followed by a beep. If you do not hear all of these sounds, discontinue use of the ventilator and have it serviced.

1. Power on the ventilator with the **ON/Shutdown** key.
2. Verify the ventilator operation, as described on page 5-8.

Shutting down the ventilator

Shut down the ventilator as follows:

1. Press and release the **ON/Shutdown** key. The **Shutdown** window opens.
2. Select **Ventilator Shutdown**. The ventilator shuts down.

![Shutdown window](image)

NOTE: Improper shutdown may cause a **Power has been restored** message the next time the ventilator is turned on.

NOTE: If the screen is blank and the dialogue box cannot be displayed, shut down the ventilator by pressing the **ON/Shutdown** key, then the Accept button on the top-right front of the ventilator.

Training

Product training is available. Contact your local Philips sales representative or Philips Customer Support for assistance. Call 1-800-225-0230 for ordering and 1-800-722-9377 for service.
General information
Chapter 4. Principles of operation

System operational overview

The Respironics V60/V60 Plus Ventilator is a microprocessor-controlled pneumatic system that delivers a mixture of air and oxygen. It is powered by AC with battery backup to protect against power failure or unstable power and to facilitate intrahospital transport. The ventilator’s pneumatics deliver gas and its electrical systems control pneumatics, monitor the patient, and distribute power.

The user provides inputs to the ventilator through a touchscreen, a key panel, a navigation ring (legacy versions), and an Accept button. These inputs become instructions for the pneumatics to deliver a precisely controlled gas mixture to the patient. Pressure and flow sensors provide feedback, which is used to adjust gas delivery to the patient. Monitored data based on sensor inputs is also displayed by the graphical user interface.

The ventilator’s gas delivery and monitoring functions are cross-checked. This cross-checking helps prevent simultaneous failure of these two main functions and minimizes the possible hazards of system failure.

A comprehensive system of visual and audible alarms helps ensure the patient’s safety. Clinical alarms can indicate an abnormal physiological condition. Technical alarms, triggered by the ventilator’s self-tests, can indicate a hardware or software failure. In the case of some technical alarms, limited ventilation is provided to give the user time for corrective actions. When a condition is critical enough to possibly compromise safe ventilation, the ventilator is placed into the ventilator inoperative state, in which oxygen flow and blower operation are disabled.

The ventilator has several means to ensure that safe patient or respiratory pressures are maintained. The maximum working pressure is ensured by the high inspiratory pressure (HIP) alarm limit. If the set high pressure limit is reached, the ventilator cycles into exhalation.
Principles of operation

Pneumatic system operation

The ventilator uses ambient air and high-pressure oxygen (Figure 4-1). Air enters through an inlet filter. Oxygen enters though a high-pressure inlet, and a proportioning valve provides the operator-set concentration. The system mixes the air and oxygen, pressurizes it in the blower, and then regulates it to the user-set pressure. To do this, the ventilator compares the proximal (patient) pressure measurement with the ventilator outlet (machine) pressure, and adjusts the machine pressure to compensate for the pressure drop across the inspiratory filter, patient circuit, and humidifier. This helps ensure accurate and responsive pressure delivery and leak compensation.

![Figure 4-1: Respironics V60/V60 Plus Ventilator gas delivery system](image)

Breath delivery characteristics

Control variable

Breaths delivered by the Respironics V60/V60 Plus Ventilator are pressure controlled. In the AVAPS mode, the ventilator’s applied pressure is automatically adjusted over a period of time to maintain a target tidal volume.

Triggering, cycling, and leak adaptation

Unlike other ventilators, the Respironics V60/V60 Plus Ventilator does not require you to set triggering and cycling sensitivity or to adjust baseline flow. The ventilator’s unique Auto-Trak Sensitivity algorithm adjusts these automatically; see “Auto-Trak Sensitivity” on page 4-3.

Baseline pressure

A positive baseline pressure (EPAP or CPAP) may be set for all breaths in all modes.
Principles of operation

Pressure rise time
The operator-set Rise Time defines the time required for inspiratory pressure to rise to the set (target) pressure.

Negative pressures
There are no negative pressures generated during exhalation.

Oxygen concentration
The Respironics V60/V60 Plus Ventilator incorporates an oxygen mixer. Oxygen concentration can be set in all modes.

Auto-Trak Sensitivity
An important characteristic of the Respironics V60/V60 Plus Ventilator is its ability to recognize and compensate for intentional and unintentional leaks in the system, and to automatically adjust its triggering and cycling algorithms to maintain optimum performance in the presence of leaks. This is called Auto-Trak Sensitivity. The following subsections describe this function in detail.

Triggering
Breaths are patient (flow) triggered in all modes, typically when patient effort causes a certain volume of gas to accumulate above baseline flow (volume method). An inspiration is also triggered when the patient inspiratory effort distorts the expiratory flow waveform sufficiently (shape signal method; see page 4-4).

Cycling
Cycling to exhalation occurs in these cases:

- Patient expiratory effort distorts the inspiratory flow waveform sufficiently (shape signal method). See “Shape signal method of cycling and triggering.” on page 4-4.
- Patient flow reaches the spontaneous exhalation threshold (SET). See “SET method of cycling.” on page 4-4.
- After 3 seconds at the IPAP level (timed backup safety mechanism)
- When a flow reversal occurs, typically due to a mask or mouth leak...
Principles of operation

**Shape signal method of cycling and triggering.** The shape signal or “shadow trigger” method uses a mathematical model derived from the flow signal. A new flow signal (shape signal) is generated by offsetting the signal from the actual flow and delaying it (Figure 4-2). This intentional delay causes the flow shape signal to be slightly behind the patient’s flow signal. If there is a sudden change in patient flow, the patient’s flow signal crosses the shape signal; this results in a trigger or a cycle. As a result, a sudden decrease in expiratory flow from an inspiratory effort will cross the shape signal and create a signal for ventilator triggering.

![Figure 4-2: Shape signal](image)

**SET method of cycling.** Patient flow reaches the spontaneous exhalation threshold (SET); see Figure 4-3. The SET represents the intersection of the flow waveform and a line of a given slope. SET is updated each breath.

![Figure 4-3: Spontaneous exhalation threshold (SET)](image)
Principles of operation

Leak adaptation
Noninvasive ventilation in particular may involve considerable leakage around the mask or through the mouth. Some leakage is known or intentional: it is a characteristic of the mask/patient interface design. So that it can accurately adjust its baseline flow, the ventilator has you enter the intentional leakage value specific to the mask/patient interface (“Selecting the mask and exhalation port” on page 6-12). Other leakage is unpredictable or unintentional, and it changes as the patient’s breathing pattern changes.

To maintain prescribed pressures in the presence of leakage, the ventilator adjusts its baseline flow. Because the unintentional part of the leakage may constantly change, the ventilator recalculates the baseline flow each breath at the end of exhalation. The ventilator uses two main mechanisms to update its baseline flow: expiratory flow adjustment and tidal volume adjustment.

Expiratory flow adjustment. Every breath, at end-exhalation, the ventilator updates its flow baseline. At end-exhalation patient flow is assumed to be zero, so any difference between actual patient flow and the original baseline flow indicates a change in leakage. Figure 4-4 shows how the ventilator adjusts the baseline.

Figure 4-4: Expiratory flow adjustment
Principles of operation

**Tidal volume adjustment.** Every breath, the ventilator compares the inspiratory and expiratory tidal volumes. Any difference is assumed to be due to an unintentional circuit leak. The ventilator adjusts the baseline to reduce this tidal volume difference for the next breath. Figure 4-5 shows how the ventilator adjusts the baseline.

![Figure 4-5: Tidal volume adjustment](image)

---

**Auto-Trak+ (optional)**

The Auto-Trak+ option for the Respironics V60/V60 Plus Ventilator lets you further adjust the level of Auto-Trak Sensitivity, a feature that recognizes and compensates for intentional and unintentional leaks. This algorithm has multiple breath trigger and cycle thresholds. When you adjust Auto-Trak+ settings, you adjust these multiple trigger and/or cycle thresholds simultaneously, retaining all the auto-adaptive features of Auto-Trak Sensitivity.

The Normal Auto-Trak settings work well for most patients. Pediatric patients, however, may benefit from more sensitive trigger settings, while some adult patients may benefit from more or less sensitive cycle settings.

---

**High flow therapy**

High flow therapy (HFT) enables delivery of a humidified gas mixture at an operator-set flow rate via a high flow nasal cannula interface or tracheal adapter. The principle mechanism of action for high flow therapy is delivering a known FiO₂ at a flow rate equal to or greater than the patient's peak flow, thus minimizing dilution of the gas.

HFT provides blended gas to the patient at a targeted flow. Both O₂ concentration and flow are set by the clinician. Heated humidification is recommended during high flow therapy.
Principles of operation

HFT controls flow instead of pressure and is accessed only while in Standby mode. Patient alarms are not available during high flow therapy. This therapy is not considered a breath delivery mode.

HFT requires 3.00 software and above.

Ventilation modes

The Respironics V60/V60 Plus Ventilator operates in the following ventilation modes:

- CPAP (continuous positive airway pressure) mode
- S/T (spontaneous/timed) mode
- PCV (pressure-controlled ventilation) mode
- AVAPS (average volume-assured pressure support) mode
- PPV (proportional pressure ventilation) mode (optional)

Table 4-1 summarizes the characteristics of these modes. Note that on the ventilator, the Timed breath indicator means the breath is ventilator triggered, while the Spont breath indicator means the breath is patient triggered.

<table>
<thead>
<tr>
<th>Mode</th>
<th>Timed breaths</th>
<th>Spont breaths</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Trigger*</td>
<td>Limit†</td>
</tr>
<tr>
<td>CPAP</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>PCV</td>
<td>Time</td>
<td>Pressure</td>
</tr>
<tr>
<td>S/T</td>
<td>Time</td>
<td>Pressure</td>
</tr>
<tr>
<td>AVAPS</td>
<td>Time</td>
<td>Pressure</td>
</tr>
<tr>
<td>PPV</td>
<td>Time</td>
<td>Pressure</td>
</tr>
</tbody>
</table>

* A trigger variable starts inspiration.
† A limit variable can reach and maintain a preset level before inspiration ends but it does not end inspiration.
‡ A cycle variable is a measured parameter used to end inspiration.
**Principles of operation**

**CPAP mode**

In the CPAP (continuous positive airway pressure) mode, the ventilator functions as a demand flow system, with the patient triggering all breaths and determining their timing and size. The patient triggers and cycles based on the ventilator’s Auto-Trak Sensitivity algorithms. The control settings active in the CPAP mode are shown in Figure 4-6. Figure 4-7 shows CPAP mode waveforms.

The C-Flex feature setting enhances traditional CPAP by reducing the pressure at the beginning of exhalation – a time when patients may be uncomfortable with CPAP – and returning it to the set CPAP level before the end of exhalation.

![Figure 4-6: CPAP controls](image)

![Figure 4-7: CPAP waveforms](image)
PCV mode
The PCV (pressure-controlled ventilation) mode delivers pressure-controlled breaths, either triggered by the ventilator (Timed) or the patient (Spont). The control settings active in the PCV mode are shown in Figure 4-8. The IPAP setting defines the applied inspiratory pressure for all breaths. If the patient fails to trigger a breath through Auto-Trak within the interval determined by the rate setting, the ventilator triggers a mandatory breath. The I-Time setting is the cycle criterion for all breaths. Figure 4-9 shows a PCV mode pressure waveform.

**Figure 4-8: PCV controls**

- **IPAP**: 12 cmH2O
- **Rate**: 4 BPM
- **EPAP**: 4 cmH2O
- **I-Time**: 1.00 sec
- **Rise**: 2
- **O2**: 21%
- **I-Time**: 1/Rate
- **Machine-triggered (Timed) breath**
- **Patient-triggered (Spont) breath**

**Figure 4-9: PCV pressure waveform**
Principles of operation

S/T mode
The S/T (spontaneous/timed) mode guarantees breath delivery at the user-set rate. It delivers pressure-controlled, time-cycled mandatory and pressure-supported spontaneous breaths, all at the IPAP pressure level. If the patient fails to trigger a breath within the interval determined by the Rate setting, the ventilator triggers a mandatory breath with the set I-Time. The patient triggers and cycles based on the ventilator’s Auto-Trak Sensitivity algorithms. The control settings active in the S/T mode are shown in Figure 4-10. Figure 4-11 shows an S/T mode pressure waveform.

![Active Mode: S/T](image)

**Figure 4-10: S/T controls**

![Pressure waveform](image)

**Figure 4-11: S/T pressure waveform**
AVAPS mode

NOTE: When you adjust AVAPS minimum and maximum pressures, remember that IPAP is adjusted to meet the target value. If the calculated target pressure is outside of the minimum and maximum pressure range, the target volume will not be achieved.

Unlike most pressure modes, the AVAPS (average volume-assured pressure support) mode delivers a target tidal volume. It achieves the target volume by regulating the pressure applied following an initial pressure ramp-up. The AVAPS mode delivers time-cycled mandatory breaths and pressure-supported spontaneous breaths.

If the patient fails to trigger a breath within the interval determined by the Rate control, the ventilator triggers a mandatory breath with the set I-Time. Mandatory and spontaneous breaths are delivered at a pressure that is continually adjusted over a period of time to achieve the volume target, V_T. Min P and Max P define the minimum and maximum pressures that can be applied. The patient triggers and cycles based on the ventilator’s Auto-Trak Sensitivity algorithms.

At start-up, AVAPS applies an inspiratory pressure equal to one of the following, whichever is greater:

- EPAP + (target volume / 60 ml/cmH2O)
- EPAP + 8 cmH2O
- Min P

The control settings active in the AVAPS mode are shown in Figure 4-12. Figure 4-13 shows AVAPS mode waveforms.

![Figure 4-12: AVAPS controls](image-url)
Principles of operation

Figure 4-13: AVAPS waveforms
**PPV mode (optional)**

The PPV (proportional pressure ventilation) mode provides patient-triggered breaths that deliver pressure in proportion to patient effort. Additionally a user-settable backup rate activates machine-triggered, pressure-limited, and time-cycled breaths in the case of apnea. In the PPV mode, patient effort determines the pressure, flow, and tidal volume delivered by the ventilator. The ventilator responds to patient effort, allowing the patient to determine when to start and end a breath. Additionally flow and pressure change based on the patient’s efforts throughout inspiration.

**The physics behind PPV.** Two forces oppose ventilation, resistance and elastance.

Resistance is the impedance to air movement in the airways:

\[
\text{Pressure/Flow} = \text{Resistance}
\]

Airway resistance in healthy adults ranges from approximately 0.5 to 2.5 cmH₂O/L/s.

Elastance is the elastic opposition to ventilation or the tendency of the lungs to resist inflation (elastance is the reciprocal of compliance):

\[
\text{Pressure/Volume} = \frac{1}{\text{Compliance}} = \text{Elastance}
\]

The compliance of lungs and chest wall for a healthy adult is approximately 0.1 L/cmH₂O, resulting in an elastance value of 10 cmH₂O/L.

The inspiratory muscles, therefore, must generate force to overcome the resistance and elastance of the respiratory system. The proximal airway pressure is the net result of this contraction of these muscles: it is the force of the inspiratory muscle contraction minus both the pressure needed to generate airflow (overcome respiratory system resistance) and the pressure generated to inflate the lungs (overcome respiratory system elastance).

**How PPV works.** The delivery of a PPV breath is controlled by the maximum elastance (volume) assist (Max E), maximum resistance (flow) assist (Max R), and PPV % settings. The actual delivered assistance to overcome elastance is the product of PPV % and Max E. The actual delivered assistance to overcome resistance is the product of PPV % and Max R. In general, Max E should be set relative to the respiratory elastance and Max R should be set relative to the respiratory resistance, although you do not need to know the actual value of either to apply PPV. You adjust assist levels to optimize patient comfort. The resultant pressure support delivered in the PPV mode is the resistance assist times patient flow plus the elastance assist times the patient volume. Because the patient completely controls ventilatory output, PPV may significantly improve patient-ventilator synchrony and ultimately, patient comfort.

The PPV backup rate ensures that the patient receives a minimum number of breaths per minute if the spontaneous breathing rate falls below the Rate

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Principles of operation

setting. If the patient fails to trigger a breath within the interval determined by the Rate control, the ventilator triggers a Timed (backup) breath with the set I-Time, Rise, and IPAP settings.

The control settings active in the PPV mode are shown in Figure 4-14.

Figure 4-14: PPV controls

Figure 4-15 shows PPV mode waveforms. Note how volume and pressure increase as does the ventilatory demand of the patient. Max V (PPV maximum volume limit) and Max P (PPV maximum pressure limit) are used to prevent the delivery of excessive pressure or volume. More information about these limits is provided in “About Max V and Max P alarms and alarm limits” on page 6-7.

Figure 4-15: PPV waveforms
Principles of operation

Oxygen mixing

The ventilator’s oxygen mixer regulates and proportions oxygen into the air from the blower according to the $O_2$ setting. The delivered oxygen accuracy is ±5% of the set value up to the maximum oxygen flow available. The ventilator can deliver up to 240 L/min of air/oxygen mix to assist in managing uncontrolled leaks during noninvasive ventilation.

Many hospital oxygen supply systems, however, cannot meet such high flow demands. Under extraordinary conditions (high $O_2$ setting plus high leak, and/or high patient demand) where demand exceeds available oxygen system flow, the ventilator provides additional air flow from the blower to ensure the target pressure is met. Under such conditions, the accuracy of delivered oxygen may be affected. Figure 4-16 shows the effect on the delivered oxygen concentration as the maximum oxygen system flow is exceeded. This graph assumes a continuous flow demand. Normally the higher “peak” flow is only needed during inspiration, so this is a worst case scenario.

Assumptions: At an $O_2$ setting of 100% and an oxygen supply with a 50 psig inlet pressure capable of delivering up to 160 L/min.

Figure 4-16: $O_2$ concentration as a function of total ventilator flow
Principles of operation
Chapter 5. Setting up the ventilator for use

Set up the ventilator for each patient use as described in this chapter. For first-time installation, refer to Appendix A. For use with high flow therapy (HFT), set up the ventilator as described in this chapter, then refer to Chapter 7, High flow therapy.

Connecting oxygen

**WARNING:** Connect the ventilator only to an appropriate medical-grade oxygen source.

**WARNING:** To ensure accuracy of oxygen administration and to monitor for the presence of contamination (incorrect gas connected), use an external oxygen monitor to verify the oxygen concentration in the delivered gas.

**WARNING:** To reduce the risk of fire, do not use a high-pressure oxygen hose that is worn or contaminated with combustible materials like grease or oil.

**WARNING:** To reduce the risk of hypoxia, connect only oxygen to the high-pressure connector at the rear of the ventilator.

**WARNING:** To reduce patient risk of oxygen toxicity, keep free-flowing oxygen away from air inlet of ventilator.

**CAUTION:** To prevent possible damage to the ventilator, ensure that the connection to the oxygen supply is clean and unlubricated, and that there is no water in the oxygen supply gas.

Connect the oxygen hose to an appropriate high-pressure oxygen source using country-specific O₂ connectors, as applicable.

Use of SIS connectors and supplemental oxygen accessories such as the O₂ manifold requires higher oxygen supply pressures. Consult Table 11-9 on page 11-7 for appropriate oxygen pressure ranges.

Installing an oxygen analyzer/monitor

Install an Analytical Industries 2000M oxygen analyzer/monitor, and follow the manufacturer’s instructions for setup, alarms, and calibration.

The Analytical Industries 2000M monitor includes user-settable high and low oxygen % alarms and is approved for use with the V60/V60 Plus Ventilator. Refer to the monitor instructions for use for detailed instructions on the proper setup and operation of the oxygen monitor.

**NOTE:** The V60/V60 Plus Ventilator also incorporates a loss of oxygen supply alarm to further protect the patient from low oxygen supply pressure conditions.
Setting up the ventilator for use

### Connecting to AC power

| WARNING: | To reduce the risk of electric shock, connect the ventilator to an AC supply mains with protective earth only. |
| WARNING: | Do not use extension cords, adapters, or power cords with the ventilator that are not approved by Respironics. |
| WARNING: | To prevent unintentional disconnection of the power cord, always use the correct, Philips-supplied power cord and lock it into place with the power cord retainer before you switch the ventilator on. The retainer is designed to hold the connector end of the Philips-supplied cord securely in place. |
| WARNING: | To reduce the risk of electric shock, regularly inspect the AC power cord and verify that it is not frayed or cracked. |
| WARNING: | To reduce the risk of strangulation, route the power cord to avoid entanglement. |
| CAUTION: | For 120 V equipment, grounding reliability can only be achieved when it is connected to an equivalent receptacle marked “hospital only” or “hospital grade.” |

Plug the power cord into a grounded outlet that supplies AC power between 100 and 240 V, 50/60 Hz.

Always check the reliability of the AC outlet. If you are using a 120 V outlet, make sure that it is hospital grade.
Installing the patient circuit

WARNING: To reduce the risk of strangulation from patient tubing, use a tubing support arm and secure the proximal pressure line with clips.

WARNING: To prevent possible patient injury and possible water damage to the ventilator, make sure the humidifier is set appropriately.

WARNING: To prevent possible patient injury and equipment damage, do not turn the humidifier on until the gas flow has started and is regulated. Starting the heater or leaving it on without gas flow for prolonged periods may result in heat build-up, causing a bolus of hot air to be delivered to the patient. Circuit tubing may melt under these conditions. Turn the heater power switch off before stopping gas flow.

WARNING: To reduce the risk that the patient will aspirate condensed water from the breathing circuit, position any humidifier lower than both the ventilator and the patient.

WARNING: To reduce the risk of fire, use only patient circuits intended for use in oxygen-enriched environments. Do not use antistatic or electrically conductive tubing.

WARNING: To prevent patient or ventilator contamination, always use a main flow bacteria filter on the patient gas outlet port. Filters not approved by Respironics may degrade system performance.

WARNING: During ventilation, patient exhalate is released into room air. Use of a patient circuit with a filter on its exhalation port is recommended.

WARNING: To reduce the risk of bacterial contamination or damage, handle bacteria filters with care.

WARNING: Avoid adding any components to the patient circuit that are not absolutely necessary. Additional components installed in the patient circuit can change the pressure gradient across the ventilator breathing system, increase the dead space, and adversely affect the ventilator performance.

WARNING: Any additional accessories in the patient circuit may substantially increase flow resistance and impair ventilation.

WARNING: Avoid adding resistive circuit components on the patient side of the proximal pressure line. Such components may defeat the disconnect alarm.

WARNING: Using a jet nebulizer can cause inadvertent alarms and affect the accuracy of delivered FiO₂. To reduce patient risk, use only an approved nebulizer.

NOTE: Bacteria filter must be installed onto gas outlet.

NOTE: Resistive components can include but are not limited to HMEs, proximal flow sensors, a filter at the patient connection, or a narrow diameter circuit attached to a mask.

NOTE: Under extreme conditions and a missing, ruptured, or defective bacteria filter, the entire gas pathway can become contaminated with bodily fluids or exhaled gas.

Install the patient circuit as shown in this section. For a list of compatible parts and accessories offered by Philips, see “Parts and accessories” on page C-1.
Setting up the ventilator for use

Assemble the patient circuit, including the main flow (inspiratory) bacteria filter, proximal pressure line, oxygen sensor tee, and if desired, humidifier and nebulizer.

Figure 5-1 shows the circuit configuration for noninvasive ventilation or high flow therapy. For details on high flow therapy setup, see “High flow nasal cannula setup” on page 7-2.

Figure 5-2 and Figure 5-3 show circuit configurations for noninvasive and invasive ventilation. Follow the manufacturers’ instructions for use for the individual parts.

Installing the nebulizer
For installation of the Aerogen nebulizer, follow the manufacturer’s instructions that came with the nebulizer, or visit:

www.aerogen.com/nebulization-product-support
NOTE: This circuit setup is recommended for noninvasive ventilation. It is also recommended for high flow therapy when using the AC611 high flow nasal cannula with FEP Connector to block the exhalation port.

Figure 5-1: Patient circuit, with heated-wire and humidification for noninvasive ventilation or high flow therapy
Setting up the ventilator for use

NOTE: This circuit setup is recommended for noninvasive ventilation.

Figure 5-2: Noninvasive patient circuit, without humidification
Setting up the ventilator for use

NOTE: This circuit setup is recommended for both noninvasive and invasive ventilation. Figure 5-3 shows an invasive circuit as an example.

Connecting external devices

Connect the ventilator to a remote alarm (nurse call) device and a patient monitor or other external device, if applicable.

The Respironics V60/V60 Plus Ventilator can communicate with a Philips patient monitor using the IntelliBridge Open Interface. See “Using Philips monitors and the IntelliBridge or VueLink Open Interfaces” on page B-2. The ventilator also supports the VueLink Open Interface. VueLink has been replaced by IntelliBridge, but information is included in this manual for backwards compatibility. See “Data display” on page B-3.

For more information about connecting with non-Philips systems, contact your Philips representative.
Setting up the ventilator for use

Before placing a patient on the ventilator

**WARNING:** To ensure the ventilator’s safe operation, always verify ventilator operation as described in "Verify ventilator operation" on page 5-8 before using the ventilator on a patient. If the ventilator fails any tests, remove it from clinical use immediately. Do not use the ventilator until necessary repairs are completed and all tests have passed.

**WARNING:** To reduce the risk of power failure to the ventilator, pay close attention to the battery's charge level. The battery's operation time is approximate and is affected by ventilator settings, discharge and recharge cycles, battery age, and ambient temperature.

**NOTE:** If the ventilator has a backup battery, the battery must be adequately charged to verify operation. Recharge as necessary before verifying operation. Based on the age and state of the battery, it may take up to 16 hours or more to fully charge.

**NOTE:** The backup batteries are intended for short-term use only. They are not intended to be a primary power source.

**NOTE:** We recommend that the ventilator’s batteries be fully charged before you ventilate a patient. If the batteries are not fully charged and AC power fails, always pay close attention to the level of battery charge.

Verify ventilator operation

1. Ensure that the ventilator is connected to AC power.

2. Power on the ventilator. The ventilator automatically runs a test of the backup audible alarm followed by the primary audible alarm. Verify that you hear a high-pitched tone, followed by a beep.

3. Create a patient alarm, such as a disconnect alarm.
   a. **VERIFY** that the proper alarm is annunciated (audio, visual, flashing, and alarm LED).
   b. **VERIFY** that the audio volume is adequate for the environment in which it will be used.
   c. **VERIFY** remote alarm setup, if applicable.

4. Resolve the alarm condition and manually reset the alarm.

5. Disconnect the ventilator from AC power while the ventilator is running and verify the following:
   a. **VERIFY** that the ventilator switches over to battery power (battery symbol in lower-right corner of screen is displayed).

   ![2:00 Battery Symbol]

   b. **VERIFY** that the audible alarm sounds intermittently.
   c. **VERIFY** that the yellow “Running on Internal Battery” alarm is displayed, and manually reset it.
Setting up the ventilator for use

d. VERIFY that the blue “Running on Internal Battery” message is displayed.

6. Install a fully-charged battery and reconnect the ventilator to AC power.

7. Repeat step 5 to verify that the ventilator switches over to battery power.

Running alarm tests

The ventilator performs a self-check during start-up and continuously during operation. Alarm functionality is verified by this self-check. You may also want to run alarm tests, which demonstrate the alarms' operation. Follow these steps to perform the tests.

WARNING: To prevent possible patient injury, always return alarm settings to hospital-standard values after verifying ventilator operation.

Preparation

1. Set the ventilator up as for normal ventilation, complete with breathing circuit (PN 582073 or the equivalent) and a 1-liter test lung assembly (PN 1021671).

2. Set the mode to S/T and make the following control settings: Rate: 4 BPM, IPAP: 10 cmH₂O, EPAP: 6 cmH₂O, I-Time: 1 sec, Rise: 1, Ramp: Off, O₂: 21%.


High Inspiratory Pressure

1. Lower the HIP alarm limit to 8 cmH₂O.

2. VERIFY that the High Inspiratory Pressure alarm is activated, the ventilator cycles into exhalation, and pressure falls to 6 cmH₂O (the EPAP level).

3. Raise the HIP alarm limit to 15 cmH₂O.
Setting up the ventilator for use

Low Tidal Volume
1. Raise the Lo VT alarm setting above the displayed, measured VT.
2. VERIFY that the Low Tidal Volume alarm is activated.
3. Turn the Lo VT alarm setting OFF.
4. VERIFY that the alarm resets.

Patient Disconnect
1. Disconnect the test lung.
2. VERIFY that the Patient Disconnect alarm is activated.
3. Reconnect the test lung.
4. VERIFY that the alarm resets and that the ventilator automatically resumes ventilation.

Patient Circuit Occluded
1. Disconnect the patient circuit (including bacteria filter) from the ventilator outlet, and block the ventilator outlet.
2. VERIFY that the Patient Circuit Occluded alarm is activated.
3. Unblock the outlet, and reconnect the circuit.
4. VERIFY that the alarm resets.
Using the ventilator for intra-hospital transport

**WARNING:** Always check the status of the oxygen cylinders before using the ventilator during transport.

**WARNING:** To reduce the risk of power failure to the ventilator, pay close attention to the battery’s charge level. The battery’s operation time is approximate and is affected by ventilator settings, discharge and recharge cycles, battery age, and ambient temperature.

**WARNING:** The V60/V60 Plus Ventilator requires a pressurized oxygen supply that provides a minimum flow of 175 SLPM. Do not use any devices such as valves, hoses, Grab n’ Go regulators or other brands of combined cylinder/regulators that limit supply of oxygen flow below 175 SLPM.

**WARNING:** Do not leave the ventilator unattended when stationed on an incline.

Do the following to conserve oxygen during transport with the ventilator.

- Make sure all cylinders are full (13,790 kPa/2000 psig or more).
- Make sure the cylinder regulators are turned off while the ventilator is connected to wall oxygen.
- Never turn the cylinder regulator on until you are ready to begin transport.
- Only turn one cylinder regulator on at a time. If you turn on both cylinders, they may become depleted simultaneously, leaving you with no backup oxygen.
- Whenever possible, reduce the $O_2$ setting before transport.
- Minimize all inadvertent leaks. Tighten masks prior to transport, and loosen up when patient is back on wall oxygen.
- Avoid using masks that have an exhalation port built into the mask when there is already an exhalation port in the circuit.
- Be aware that oxygen is more rapidly depleted at higher leak rates (see Figure 5-4).
Setting up the ventilator for use

Figure 5-4: Duration of cylinder oxygen (13,790 kPa/2000 psig) at various leak rates

a. $V_t = 500$ mL, Rate = 40 BPM, EPAP = 6 cmH$_2$O, IPAP = 18 cmH$_2$O

b. $V_t = 500$ mL, Rate = 20 BPM, EPAP = 6 cmH$_2$O, IPAP = 18 cmH$_2$O

Figure 5-4: Duration of cylinder oxygen (13,790 kPa/2000 psig) at various leak rates
Storing the ventilator between patient use

See “Storage between patient use” on page 10-8 for information about storing the ventilator.

MRI safety information

WARNING: The V60/V60 Plus Ventilator is MR Unsafe. Keep it outside the MRI scan room (Zone IV). It represents a projectile hazard.

Security and Privacy Information

To develop a security strategy related to the use of the ventilator, refer to the Hospital Respiratory Care product security guide. Download it from: www.philips.com/hrcmanuals.
Setting up the ventilator for use
Chapter 6. Operation

WARNING: To ensure the ventilator’s safe operation, always verify ventilator operation as described in “Verify ventilator operation” on page 5-8 before using the ventilator on a patient. If the ventilator fails any tests, remove it from clinical use immediately. Do not use the ventilator until necessary repairs are completed and all tests have passed.

NOTE: Before operation, prepare the ventilator as instructed in Chapter 5.

NOTE: Any mode and settings can be used for closed suctioning.

After power-on, the ventilator starts up in the mode and with the settings that were active before last power down. Check these settings and adjust as required. You must be familiar with using the touchscreen and navigation ring (legacy versions) to select, adjust, activate, and confirm parameters. For details, see “Before placing a patient on the ventilator” on page 5-8.

Access the ventilator setting windows from the tabs at the bottom of the screen.
Operation

Changing the mode

The active ventilation mode is displayed in the bottom, left-hand corner of the screen. Change the mode as follows. For details on modes, see “Ventilation modes” on page 4-7.

1. Open the Modes window.
2. Select the desired mode.
3. Adjust settings as desired (see “Changing individual ventilator settings” on page 6-4). Newly adjusted setting values are shown in yellow.
4. Select Activate Mode to apply.
Changing control settings

Table 6-3 on page 6-23 is an alphabetical list of the control settings with their ranges. Table 11-2 on page 11-2 shows the control settings applicable to the different modes. For more information on control settings as they apply in the different ventilation modes, see “Ventilation modes” on page 4-7.

Making batch setting changes

The ventilator allows you to make multiple changes (a “batch” of changes) at once.

**NOTE:** During a batch setting change, you cannot change the Ramp Time setting when a ramp is active.

This process applies to ventilation settings only, not to alarm settings.

1. Open the Modes window.
2. Select the active mode.
3. Adjust settings as desired (see “Changing individual ventilator settings” on page 6-4). Newly adjusted setting values are shown in yellow.
4. Select Activate Batch Change to apply.
Changing individual ventilator settings
You can make ventilator settings from the Settings window.

1. Open the Settings window.

2. Select the desired setting. As an example we will show the IPAP adjustment.

3. The setting window opens. Adjust the setting. Select Accept to apply.
Using the Ramp Time function

The Ramp Time function helps your patient adapt to ventilation by gradually increasing inspiratory and expiratory pressure (IPAP and EPAP/CPAP) from subtherapeutic to user-set pressures over a user-set interval. Table 6-3 on page 6-23 describes this function’s principles of operation.

Follow these instructions to use the Ramp Time function:

1. Select the Ramp Time button in the Settings window.

   ![Ramp Time button graphic]

   The ramp starts. As the ramp progresses, the Ramp Time button graphic fills in.

2. To change the ramp interval or to end the ramp, select the Ramp Time button again. The Ramp in Progress window opens.

   ![Ramp in Progress window]

   3. To end the ramp and apply the full IPAP and EPAP/CPAP immediately, select End Ramp.

   4. To end the ramp and start a new one, select Start New Ramp. The Ramp Time setting window opens again so that you can set up a new ramp.
Operation

Using the 100% O₂ function

NOTE: The 100% O₂ feature is available in Revision 2.30 software and above.

The 100% O₂ function delivers 100% oxygen to the patient. It is available during Screen Lock status.

Follow these instructions to use the 100% O₂ function:

1. Select the 100% O₂ button in the main GUI window.

![100% O₂ button](image)

2. The ventilator delivers 100% oxygen for two minutes. A countdown timer displays.

![Countdown timer](image)

While 100% oxygen delivery is active, you can press the +2:00 button to add two minutes more. Press Cancel to stop.
Using PPV

Follow these instructions to set up the ventilator in the PPV mode, referring to Figure 6-3. For principles of operation, see “PPV mode (optional)” on page 4-13.

1. Open the PPV Settings window.

2. Set EPAP, O2, alarm limits, and backup settings to appropriate values. The HIP alarm limit should be greater than Max P. See “Principles of operation” on page 4-1 for a detailed explanation of these settings.

3. Set the Max V and Max P limits.

4. Set alarm limits to appropriate values. The HIP alarm limit should be greater than the Max P.

About Max V and Max P alarms and alarm limits
Max V (PPV maximum volume limit) and Max P (PPV maximum pressure limit) are used to prevent the delivery of excessive pressure or volume.

WARNING: PPV limits are not intended to be the primary ventilator alarms and should not be substituted for the alarms found in the Alarm Settings window.

WARNING: To prevent the delivery of excessive pressure or volume, set the PPV limits appropriately. Delivery of excessive pressure or volume can occur from a sudden increase in mask leak, inappropriate settings, or a plugged or kinked proximal pressure line. Conversely, insufficient treatment may result if limits are set too low.

When the Max V (PPV maximum volume limit) is reached, the breath is terminated and a message is displayed. After the limit is reached in three consecutive breaths, the audible alarm sounds. A PPV waveform with Max V is shown in Figure 6-1.
When the Max P (PPV maximum pressure limit) is reached, pressure is limited but the breath is not terminated, and a message is displayed. After the limit is reached in three consecutive breaths, the audible alarm sounds. A PPV waveform with Max P is shown in Figure 6-2.

Frequent annunciation of one or both alarms typically indicates improved patient status. It may, however, indicate that the patient is more actively breathing, possibly due to agitation or a change in the patient’s level of sedation. It may also indicate an increase in leakage.

The $V_T$ (estimated exhaled tidal volume) measurement may remain below the set Max V limit even though the inspired volume exceeds Max V. This results from variable leakage, which reduces the exhaled volume in relation to the inspired volume.
Guidelines for using PPV

NOTE: The guidelines below are based on recommendations by clinicians. They do not replace the clinical judgment of a physician and should not, on their own, be used for clinical decision making.

Determining Max R and Max E settings
It is recommended you set Max R (flow assist) and Max E (volume assist) to initial values and then titrate them based on the patient’s disease process:

- Obstructive disease (COPD, asthma): Focus on Max R. Overcoming increased resistance is typically the emphasis, not volume delivery.
- Restrictive disease (neuromuscular, chest-wall deformities, obesity hypoventilation): Focus on Max E. Maintaining sufficient volume is typically the emphasis, not overcoming increased resistance.
- Mixed disease processes affecting both resistance and elastance: Titrate both Max R and Max E settings.

Suggested titration procedure Follow this procedure to titrate settings to optimize patient comfort while avoiding overassisting. See also the flow chart in Figure 6-3.

NOTE: You may also need to adjust PPV % according to patient response, as you do for the other PPV settings described below. Mask leakage, especially a sudden increase, is interpreted as patient effort by the ventilator and assisted accordingly; this may necessitate lowering the PPV % setting. However, the best solution is to maintain a minimal leak.

1. Set EPAP, O₂, alarm limits, and backup settings to appropriate values. The HIP alarm limit should be greater than Max P.

Suggested starting settings:

- EPAP 4 cmH₂O
- O₂ Current setting or per prescription
- Max P 25 cmH₂O
- Max V 1000 to 1500 mL
- PPV % 80 to 100%
- Max E 5 cmH₂O/L
- Max R 2 cmH₂O/L/s
- All other backup settings and alarms Per usual protocol

* Consider higher EPAP settings for COPD patients to treat autoPEEP as evidenced by missed triggers
2. Adjust Max E:
   a. Evaluate the patient. Check whether any of these conditions is true:
      • The patient says they are getting too much air, pressure, or volume
      • The patient is using accessory muscles to actively stop inspiration
      • The Max V or Max P limit is reached
      • The mask leak has suddenly increased
   b. If none is true, increase Max E in increments of 2 cmH₂O/L while continuing to evaluate the patient’s response.
   c. If any is true, decrease Max E by 2 cmH₂O/L, and re-evaluate. Repeat to optimize patient comfort.

3. Repeat the process above adjusting Max R, increasing and decreasing in increments of 1 cmH₂O/L/s to optimize patient comfort.

4. Repeat adjustment for Max E as needed.

5. Adjust PPV % downward as tolerated.
Figure 6-3: PPV initial setup

Start

Make initial settings:
- **EPAP**: 4 cmH₂O
- **O₂**: Current setting or per prescription
- **Max P**: 25 cmH₂O
- **Max V**: 1000 to 1500 mL
- **PPV %**: 80 to 100%
- **Max E**: 5 cmH₂O/L
- **Max R**: 2 cmH₂O/L/s
- **HIP**: > Max P
- Other settings: Per usual protocol

Max E titration

- Does the patient say they are getting too much air, pressure, or volume?
  - Yes
  - No
- Is the patient using accessory muscles to actively stop inspiration?
  - Yes
  - No
- Was the Max V or Max P limit reached?
  - Yes
  - No
- Has the mask leak suddenly increased?
  - Yes
  - No
- Increase Max E by 2 cmH₂O/L
- Reduce Max E by 2 cmH₂O/L for patient comfort

Max R titration

- Does the patient say the air is coming too fast?
  - Yes
  - No
- Is the patient using accessory muscles to actively stop inspiration?
  - Yes
  - No
- Was the Max V or Max P limit reached?
  - Yes
  - No
- Has the mask leak suddenly increased?
  - Yes
  - No
- Increase Max R by 1 cmH₂O/L/s
- Reduce Max R by 1 cmH₂O/L/s for patient comfort
- Repeat Max E titration as needed

Titrate PPV % downward as tolerated

End
**Operation**

**Changing alarm settings**

---

**WARNING:** To prevent possible patient injury, avoid setting alarm limits to extreme values, which can render the alarm system useless.

---

Some ventilator alarm settings are operator adjustable. You can adjust these at any time. Table 6-4 on page 6-26 lists the alarm settings and their ranges.

Review and adjust the alarm settings as follows:

1. Open the Alarm Settings window.

![Alarm Settings](image1)

2. Select the desired setting, adjust it, and select Accept to apply.

The ventilator annunciates an alarm when a monitored value goes out of the range bounded by the alarm limits.

---

**Selecting the mask and exhalation port**

To be able to display full leakage data plus accurate tidal and minute volumes, the ventilator must know the intentional leak characteristics of the specific mask/patient interface and exhalation port.

After power-on, the Messages list displays the current mask and port settings for 5 minutes.

![Messages](image2)

Change these settings as follows:

1. Open the Menu window.

![Menu](image3)

2. Select Mask/Port.
3. Select the desired mask/patient interface type (Table 6-1). Select Accept to apply.

![Mask/ET Selection Diagram]

For information concerning mask/port leak characteristics, see the instructions provided with each mask/port. See Appendix C for a list of masks, circuits, and related components used with the ventilator.

**Table 6-1: Mask/patient interface selections**

<table>
<thead>
<tr>
<th><strong>Mask/patient interface type</strong></th>
<th><strong>Description</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>ET/Trach</td>
<td>ET or tracheostomy tube</td>
</tr>
</tbody>
</table>
| Leak 1                          | Mask with minimal intentional leak characteristics. Enter Leak 1 for any of these Philips Respironics masks:  
  - Contour Deluxe nasal mask  
  - PerformaTrak mask  
  - AF531, AF541 (EE) |
| Leak 2                          | Mask with medium intentional leak characteristics. Enter Leak 2 for this mask:  
  - Philips Respironics PerforMax oro-nasal mask [EE]  
  - AF531, AF541 (EE) |
| Leak 3                          | AP111           |
| Leak 4                          | Reserved for future use |
| Other                           | Mask not manufactured by Philips Respironics |

* A leak symbol is printed on Respironics masks.
4. Select the desired exhalation port type (Table 6-2). Select Accept to apply.

If you select an exhalation port that is not compatible with the selected mask, Not allowed with current mask is displayed.

NOTE: In ventilation modes, ET/tracheostomy tubes and most Philips Respironics masks require the use of an exhalation port. If you selected ET/Trach or Leak 1 as a mask/patient interface, you may not select None as an exhalation port.
5. Run the exhalation port test if indicated in the table (see “Running the exhalation port test” on page 6-16 for instructions).

CAUTION: If you selected PEV or Other as an exhalation port, you must run an exhalation port test.

NOTE: If the exhalation port test is not run or if it fails, the intentional leak is unknown. Tot.Leak rather than Pt. Leak is displayed in the patient data window.
Operation

Running the exhalation port test

The exhalation port test is required and its window is automatically displayed when PEV or Other is selected.

Procedure
Run the test as follows:

1. Disconnect the patient circuit from the mask/patient interface.

2. Occlude the circuit outlet. Select Start Test.

3. Wait while the test runs.

4. Verify that Test Passed is displayed.
5. Reconnect the patient circuit to the mask/interface.

6. Select Start Ventilation to initiate ventilation.

Troubleshooting
If Test Failed is displayed, check for leaks in the patient circuit, and install an exhalation device with lower leak characteristics. Repeat test. If the exhalation port test fails again, the intentional leak is unknown and Tot.Leak rather than Pt. Leak is displayed in the patient data window.

Other functions: the Menu window

From the Menu window you can adjust user preferences.

Brightness
Use Brightness to adjust the screen for optimum daytime or nighttime viewing.

Loudness

**WARNING:** Set the alarm loudness above the ambient level. Setting the alarm loudness too low may prevent recognition of alarm conditions.

**WARNING:** Avoid blocking the alarm speakers beneath the ventilator.

Use Loudness to adjust the volume of the alarm and touchscreen audible feedback. You will hear audible feedback as you go through the selections.

The Alarm Volume Escalation status is also displayed on this screen. See “Alarm Volume Escalation” on page E-11 for more information.
Operation

Mask/Port
See “Selecting the mask and exhalation port” on page 6-12.

Vent Info (ventilator information)
The Ventilator Information window displays software version and other information specific to your ventilator.

Screen Lock
Screen Lock deactivates all buttons and tabs on the touchscreen except Alarm Silence, Alarm Reset, the Alarm/Message button, and Help. Tabs are grayed out as in this example.

This message bar is displayed at the top of the screen:

To unlock the screen, press the Accept button on the top-right front of the ventilator.

NOTE: If Screen Lock is active, the touchscreen remains locked even if an alarm becomes active.

Auto-Trak+
The Normal Auto-Trak settings work well for most patients. Pediatric patients, however, may benefit from more sensitive trigger settings, while some adult patients may benefit from more or less sensitive cycle settings.

Changing Auto-Trak+ settings

1. Select Auto-Trak+ from the Menu window.
2. Select the desired adjustment. As an example, the E-Cycle adjustment is shown below.

3. The setting window opens. Adjust the setting, referring to the pressure-time graphic which represents the effect on I-Time. Select Accept to apply.

When Auto-Trak+ is active (when either Trigger or E-Cycle is set to a value other than Normal), the ventilator setting window displays Auto-Trak+.

Additionally, after power-on the Messages list displays the Auto-Trak+ settings for 5 minutes.
Standby

Standby lets you safely suspend ventilation to temporarily disconnect the patient from the ventilator or to set up the ventilator before connecting the patient. Alarms are disabled during standby.

You can also change ventilator settings and most menu functions during standby. The settings changes are effective when you exit standby. Enter standby as follows:


NOTE: Remove the mask/patient interface in order to enter standby. The ventilator will not enter standby with a patient connected. If the patient is not disconnected, the ventilator continues breath delivery while waiting for the patient to be disconnected. The standby mode request cancels in 60 seconds if the patient remains connected.

NOTE: Standby mode disables alarms and should be used when the patient is disconnected.
2. Disconnect the patient from the ventilator now. The ventilator enters standby and displays the Standby screen.

3. To resume ventilation, reconnect the patient. When the ventilator senses a patient breathing effort, ventilation automatically resumes in the previous mode.

NOTE: You can also manually resume ventilation with the Restart Mode button.
Operation

Help function

Select the help button to display additional information.

Help messages are displayed:
Table 6-3: Modes and control settings with ranges

<table>
<thead>
<tr>
<th>Setting</th>
<th>Description</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Modes</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Modes</td>
<td>Ventilation mode</td>
<td>AVAPS, CPAP, S/T, PCV Optional: PPV</td>
</tr>
<tr>
<td><strong>Control settings</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C-Flex</td>
<td>Enhances traditional CPAP by reducing the pressure at the beginning of exhalation—a time when patients may be uncomfortable with CPAP—and returning it to the set CPAP pressure before the end of exhalation. The amount of pressure relief is determined by the C-Flex setting and the expiratory flow. The higher the setting number (1, 2 or 3) and the greater the expiratory flow, the greater the pressure relief (during the active part of exhalation only). Applies in CPAP mode only.</td>
<td>OFF, 1 to 3</td>
</tr>
<tr>
<td>CPAP</td>
<td>Continuous positive airway pressure. The baseline pressure applied during the expiratory phase. Applies in CPAP mode only.</td>
<td>4 to 25 cmH₂O</td>
</tr>
<tr>
<td>E-Cycle (optional)</td>
<td>Expiratory Cycle Sensitivity. Auto-Trak+ employs several algorithms to determine the point at which the ventilator cycles into exhalation. This setting adjusts all algorithms simultaneously. At the lowest setting (-2), inspiration terminates later, resulting in the longest inspiratory time. At the highest setting (+6), inspiration terminates earlier, resulting in the shortest inspiratory time. Normal is the Auto-Trak setting used when Auto-Trak+ is not enabled. Applies only when the optional Auto-Trak+ feature is installed.</td>
<td>-2, -1, Normal, +1 to +6</td>
</tr>
<tr>
<td>EPAP</td>
<td>Expiratory positive airway pressure. The application and maintenance of pressure above atmospheric at the airway throughout the expiratory phase of positive-pressure mechanical ventilation.</td>
<td>4 to 25 cmH₂O</td>
</tr>
<tr>
<td>IPAP</td>
<td>Inspiratory positive airway pressure. The application and maintenance of pressure above atmospheric at the airway throughout the inspiration phase of positive-pressure mechanical ventilation.</td>
<td>4 to 40 cmH₂O</td>
</tr>
</tbody>
</table>
### Operation

#### Table 6-3: Modes and control settings with ranges (continued)

<table>
<thead>
<tr>
<th>Setting</th>
<th>Description</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>I-Time (Inspiratory Time)</td>
<td>Time to deliver the required gas. Inverse ratio ventilation is not allowed.</td>
<td>0.30 to 3.00 secs</td>
</tr>
<tr>
<td>Max E</td>
<td>The maximum elastance (volume assist) value used by the PPV mode to overcome the elastance of the patient's lungs. See also PPV % setting. Applies in PPV mode only.</td>
<td>0 to 100 cmH₂O/L</td>
</tr>
<tr>
<td>Max P (AVAPS Maximum IPAP Pressure)</td>
<td>The maximum pressure to be applied.</td>
<td>6 to 40 cmH₂O</td>
</tr>
<tr>
<td></td>
<td><strong>NOTE:</strong> When you adjust the AVAPS minimum and maximum pressures, remember that IPAP is adjusted to meet the target value. If the calculated target pressure is outside of the minimum and maximum pressure range, the target volume will not be achieved.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Applies in AVAPS mode only.</td>
<td></td>
</tr>
<tr>
<td>Max P (PPV Maximum Pressure Limit)</td>
<td>The maximum pressure to be applied. When the limit is reached, the ventilator limits the pressure and displays a PPV Max P alarm message. If the condition persists for three consecutive PPV inspirations, an audible alarm also sounds. Applies in PPV mode only.</td>
<td>5 to 40 cmH₂O</td>
</tr>
<tr>
<td></td>
<td>WARNING: PPV limits are not intended to be the primary ventilator alarms and should not be substituted for the alarms found in the Alarm Settings window. WARNING: To prevent the delivery of excessive pressure or volume, set the PPV limits appropriately. Delivery of excessive pressure or volume can occur from a sudden increase in mask leak, inappropriate settings, or a plugged or kinked proximal pressure line. Conversely, insufficient treatment may result if limits are set too low.</td>
<td></td>
</tr>
<tr>
<td>Max R</td>
<td>The maximum resistance (flow assist) value used by the PPV mode to overcome pulmonary resistance. See also PPV % setting. Applies in PPV mode only.</td>
<td>0 to 50 cmH₂O/L/s</td>
</tr>
<tr>
<td>Max V (PPV Maximum Volume Limit)</td>
<td>The maximum volume to be delivered. When the limit is reached, the ventilator terminates the breath and displays a PPV Max V alarm message. If the condition persists for three consecutive PPV inspirations, an audible alarm also sounds. Applies in PPV mode only.</td>
<td>200 to 3500 mL</td>
</tr>
<tr>
<td></td>
<td>WARNING: PPV limits are not intended to be the primary ventilator alarms and should not be substituted for the alarms found in the Alarm Settings window. WARNING: To prevent the delivery of excessive pressure or volume, set the PPV limits appropriately. Delivery of excessive pressure or volume can occur from a sudden increase in mask leak, inappropriate settings, or a plugged or kinked proximal pressure line. Conversely, insufficient treatment may result if limits are set too low.</td>
<td></td>
</tr>
<tr>
<td>Setting</td>
<td>Description</td>
<td>Range</td>
</tr>
<tr>
<td>----------------------------------------------</td>
<td>----------------------------------------------------------------------------</td>
<td>--------------</td>
</tr>
<tr>
<td>Min P (AVAPS Minimum IPAP Pressure)</td>
<td>The minimum pressure to be applied.</td>
<td>5 to 30 cmH₂O</td>
</tr>
<tr>
<td>NOTE: When you adjust the AVAPS minimum and maximum pressures, remember that IPAP is adjusted to meet the target value. If the calculated target pressure is outside of the minimum and maximum pressure range, the target volume will not be achieved.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Applies in AVAPS mode only.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>O₂</td>
<td>Oxygen concentration to be delivered.</td>
<td>21 to 100%</td>
</tr>
<tr>
<td>PPV %</td>
<td>Percentage of PPV assist or gain. This gain is applied to the Max E and Max R settings, yielding the applied Elastance and Resistance assist values. Applies in PPV mode only.</td>
<td>0 to 100%</td>
</tr>
<tr>
<td>PPV % chart</td>
<td>Max E and Max R are multiplied by PPV % to obtain the applied Elastance assist and Resistance assist values. Here a Max R setting of 4 cmH₂O/L/s and a PPV % setting of 30% yield a Resistance assist value of 1.2 cmH₂O/L/s.</td>
<td></td>
</tr>
<tr>
<td>Ramp Time</td>
<td>An interval during which time the ventilator linearly increases pressure, helping to reduce patient anxiety. Initial CPAP/EPAP = ( \frac{CPAP/EPAP + 4 \text{ cmH}_2\text{O}}{2} ) Initial IPAP = Initial EPAP + ( \frac{(IPAP - EPAP)}{2} )</td>
<td>OFF, 5 to 45 min</td>
</tr>
<tr>
<td>Rate (Respiratory Rate)</td>
<td>Respiratory frequency or number of breaths per minute. Inverse ratio ventilation is not allowed.</td>
<td>4 to 60 BPM</td>
</tr>
<tr>
<td>Rise (Rise Time)</td>
<td>Speed with which inspiratory pressure rises to the set (target) pressure. If the Rise Time is insufficient to reach the target IPAP pressure, adjust the Rise Time or I-Time setting.</td>
<td>1 to 5 (1 is fastest)</td>
</tr>
</tbody>
</table>
### Operation

Table 6-3: Modes and control settings with ranges (continued)

<table>
<thead>
<tr>
<th>Setting</th>
<th>Description</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trigger (optional)</td>
<td>Trigger Sensitivity. Auto-Trak+ employs several algorithms to determine the point at which the inspiration begins. The larger the value, the more sensitive the trigger (that is, the patient can trigger inspiration with less effort). Normal is the Auto-Trak setting used when Auto-Trak+ is not enabled. Applies only when the optional Auto-Trak+ feature is installed.</td>
<td>Normal, +1 to +7</td>
</tr>
<tr>
<td>VT (AVAPS Target Tidal Volume)</td>
<td>Target tidal volume to be delivered during inspiration. The ventilator meets this target by adjusting the inspiratory pressure with each breath. Applies in AVAPS mode only.</td>
<td>200 to 2000 mL</td>
</tr>
</tbody>
</table>

Table 6-4 lists the settable alarms. (For a complete list of non-settable alarms, including Patient Disconnect, Occlusion, Pressure Regulation High, and other alarms, refer to Table 9-3 on page 9-7.)

Table 6-4: Alarm settings

<table>
<thead>
<tr>
<th>Setting</th>
<th>Description</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hi Rate (High Rate Alarm)</td>
<td>High total breath rate.</td>
<td>5 to 90 BPM</td>
</tr>
<tr>
<td>Lo Rate (Low Rate Alarm)</td>
<td>Low total breath rate.</td>
<td>1 to 89 BPM</td>
</tr>
<tr>
<td><strong>NOTE:</strong></td>
<td>In non-CPAP modes, the Low Rate Alarm is essentially off if set below the Respiratory Rate setting.</td>
<td></td>
</tr>
<tr>
<td>Hi VT (High Tidal Volume Alarm)</td>
<td>High exhaled tidal volume.</td>
<td>200 to 3500 mL</td>
</tr>
<tr>
<td>Lo VT (Low Tidal Volume Alarm)</td>
<td>Low exhaled tidal volume.</td>
<td>OFF to 1500 mL</td>
</tr>
<tr>
<td>HIP (High Inspiratory Pressure Alarm)</td>
<td>High pressure at the patient airway.</td>
<td>5 to 50 cmH₂O</td>
</tr>
<tr>
<td>LIP (Low Inspiratory Pressure Alarm)</td>
<td>Low pressure at the patient airway.</td>
<td>OFF to 40 cmH₂O</td>
</tr>
</tbody>
</table>
NOTE: In the S/T and PCV modes, the LIP alarm should be set 3-5 cmH₂O below the IPAP level. When set in this manner, the alarm works in conjunction with the LIP T alarm to indicate if there is a failure to trigger between the two pressure levels. It will also alert the clinician to pressure degradation due to excessive leaks. See figure below.

<table>
<thead>
<tr>
<th>Setting</th>
<th>Description</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>HIP alarm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LIP alarm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>IPAP</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LIP T (Low Inspiratory Pressure Delay Time)</td>
<td>The interval from the detection of low inspiratory pressure until the alarm becomes active.</td>
<td>5 to 60 secs</td>
</tr>
<tr>
<td>Lo ( V_e ) (Low Minute Ventilation Alarm)</td>
<td>Low expiratory minute volume.</td>
<td>OFF to 99.0 L/min</td>
</tr>
</tbody>
</table>
Operation
Chapter 7. High flow therapy

The high flow therapy (HFT) feature is available for 3.00 software and above, as well as V60 Plus. HFT is accessed from the Standby mode. For more information, see “Standby” on page 6-20.

For principles of operation, see “High flow therapy” on page 4-6.

WARNING: When transitioning from a high flow therapy interface to an NIV mask, ensure that an exhalation port is placed in the circuit and is unobstructed to reduce the risk of CO₂ rebreathing.

WARNING: When transitioning from ventilation to high flow therapy, remove the NIV mask and use only a Philips-approved high flow patient interface to minimize pressure build-up and patient discomfort.

WARNING: When transitioning from high flow therapy to ventilation, remove the high flow nasal cannula as it is restrictive and may defeat alarms such as patient disconnect. Using a high flow nasal cannula in an NIV mode may lead to hypercarbia due to the inability to provide pressure support.

WARNING: Patient alarms are not available during high flow therapy (HFT) as the therapy uses an open system. A high flow nasal cannula occupies only a portion of the nares and patients can breathe through their mouth, which prevents estimation of patient parameters such as tidal volume, respiratory rate, pressure, and minute ventilation. Provide external monitoring, including oximetry, to inform the clinician of a change in the patient’s condition.

WARNING: During high flow therapy (HFT), verify that an occlusive patient interface is not being used. Occlusive patient interfaces include a cannula fully sealed within the nares, an NIV mask, or a direct connection to a tracheostomy tube or endotracheal tube. Remove any occlusive interface immediately as this may expose the patient to unintended high pressures.

NOTE: High flow therapy (HFT) is accessed only from the Standby window. Standby mode cannot be entered if a high flow nasal cannula is connected to the circuit.

Circuit setup

See “Installing the patient circuit” on page 5-3 for circuit configuration.
High flow therapy

High flow nasal cannula setup

Use either the AC611 high flow nasal cannula with FEP Connector (Figure 7-1) or the AC611 high flow nasal cannula 22 mm (Figure 7-3), which connects directly to the patient circuit.

Using the FEP Connect for high flow therapy

NOTE: This section applies only if you are using the AC611 high flow nasal cannula with filter exhalation port (FEP) connector (“FEP Connect”) to administer high flow therapy.

High flow nasal cannula setup

1. Insert the AC611 high flow nasal cannula with FEP Connector into the FEP, making sure the perforations in the port are completely blocked.

![Figure 7-1: High flow nasal cannula using the FEP Connect](image)

NOTE: Follow your institutional guidelines for infection control and single-patient use interfaces.
Figure 7-2: Proximal pressure line anchored to circuit with tubing clips

Changing from ventilation to high flow therapy (HFT)

Follow the instructions in this chapter. Immediately after starting HFT disconnect the proximal pressure line from the ventilator.

Changing from high flow therapy (HFT) to ventilation

Follow the instructions in this chapter. Immediately before activating ventilation, reconnect the proximal pressure line to the ventilator.
High flow therapy

Connecting directly to a 22 mm circuit

Remove the FEP, and connect the high flow nasal cannula with 22 mm connector directly to the circuit.

Changing from an NIV mode to high flow therapy

Follow these instructions to use the V60/V60 Plus Ventilator for high flow therapy (HFT).

1. Select **Standby**. The Entering Standby window opens.

2. Remove the patient mask or ET interface to enter Standby.

3. Install a Philips-approved high flow nasal cannula (Figure 7-1 and Figure 7-3 above) or a high flow tracheostomy interface on the patient circuit.

4. Select **HFT**.
5. From the Active Mode window, you can adjust Flow and $O_2\%$.

6. Press **Start HFT**.

7. The High Flow Therapy Active message is displayed during HFT.

8. Apply the HFT interface to the patient.

9. Note the low priority alarm stating that patient alarms are disabled during HFT. Press alarm reset to confirm this message.

---

**Viewing and pausing the HFT graph**

A flow graph is displayed during high flow therapy. Press the Pause button to view an event.
High flow therapy

Changing from high flow therapy to an NIV mode

1. Verify that the high flow nasal cannula is removed from the patient and disconnected from patient circuit.

2. Select Standby to open the Standby window.

3. Press the Enter Standby button.

4. In the Select Therapy window, press Ventilation.

5. Replace the high flow patient interface with a Philips-approved NIV mask.


7. Install the appropriate interface on the patient.

8. Verify that the ventilator detects the patient’s breath to activate ventilation, or press the Start Mode button.

HFT alarms and messages

Table 7-1 is a list of alarms and other messages displayed by the ventilator, along with descriptions, suggested corrective actions, and other information. The ID (identifier) listed with the priority type is the priority number of the alarm. This priority number determines the order of alarm message display. Unless otherwise indicated, alarms listed.
### Table 7-1: HFT Alarm and other messages: summary and troubleshooting

<table>
<thead>
<tr>
<th>Message</th>
<th>Description</th>
<th>Corrective Action</th>
<th>Priority type (ID)</th>
<th>Manually resettable</th>
<th>Autoresettable</th>
<th>Silenceable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cannot Reach Target Flow</td>
<td>Displays when HFT (high flow therapy) is active. Indicates that flow target is not achieved.</td>
<td>Check the patient. Check that the high flow nasal cannula size is appropriate for the flow setting. Check that an occlusive interface is NOT in use (a cannula fully sealed within the nares, an NIV mask or direct connection to an ETT/trach). Check for an occlusion, kink or liquid in the patient circuit.</td>
<td>Low (66)</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Patient alarms are disabled during HFT</td>
<td>Displays when HFT (high flow therapy) is active. Patient alarms are not available in this therapy.</td>
<td>Manually reset to confirm and clear the audible alarm.</td>
<td>Low/Information (68)</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Patient Circuit Occluded</td>
<td>Displays when HFT (high flow therapy) is active. Gas flow to the patient is obstructed.</td>
<td>Check the patient. Check that an occlusive interface is NOT in use (a cannula fully sealed within the nares, an NIV mask or direct connect to an ETT/trach). Check for an occlusion, kink or liquid in the patient circuit. If problem persists, provide alternative ventilation.</td>
<td>High (67)</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Check Vent: Proximal Pressure Sensor Range Error</td>
<td>Displays when HFT (high flow therapy) is active. Proximal pressure is out of range.</td>
<td>When using a high flow nasal cannula with FEP Connect, the proximal pressure line should be disconnected from the ventilator port during HFT. If this message is seen when switching the patient from ventilation to HFT, check to make sure the proximal pressure line is disconnected. If the message still occurs, provide alternative ventilation and have the ventilator serviced.</td>
<td>High (12)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>
High flow therapy
Chapter 8. Patient monitoring

The ventilator displays numeric patient data in the patient data window and real-time graphics in the waveform window (Figure 8-1). Numeric patient data is updated every breath. Table 8-1 on page 8-2 lists the ventilator’s monitored parameters.

Display conventions

The following symbols may be displayed in place of numeric values:

*** Data is not valid, and/or ventilator is in standby mode or disconnected

+++ Data is over range

--- Data is under range
Patient monitoring

Table of monitored parameters

Table 8-1: Monitored parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Patient data window</strong></td>
<td></td>
</tr>
<tr>
<td>Breath phase/trigger indicator</td>
<td><strong>Spont</strong> (spontaneous): Inspiratory phase, patient-triggered breath (color: turquoise)</td>
</tr>
<tr>
<td></td>
<td><strong>Timed</strong>: Inspiratory phase, ventilator-triggered breath (color: orange)</td>
</tr>
<tr>
<td></td>
<td><strong>Exhale</strong>: Expiratory phase (color: blue)</td>
</tr>
<tr>
<td>PIP</td>
<td>Peak inspiratory pressure. The highest patient pressure during the previous breath cycle.</td>
</tr>
<tr>
<td>Pt. Leak</td>
<td>Estimated patient leak or unintentional leak. Average during the previous breath cycle. Displayed only after a suitable exhalation port and mask/patient interface are selected.</td>
</tr>
<tr>
<td>Pt. Trig</td>
<td>Patient-triggered breaths, as a percentage of total breaths over the last 15 minutes.</td>
</tr>
<tr>
<td>Rate</td>
<td>Respiratory rate or total breathing frequency. Moving average over the last 6 breaths (or 15 seconds).</td>
</tr>
<tr>
<td>$T_I/T_{TOT}$</td>
<td>Inspiratory duty cycle or inspiration time divided by total cycle time. Moving average over the last 8 breaths.</td>
</tr>
<tr>
<td>Tot.Leak</td>
<td>Estimated total leak. Average during the previous breath cycle. Displayed before a suitable exhalation port and mask/patient interface are selected.</td>
</tr>
<tr>
<td>$V_E$</td>
<td>Estimated minute ventilation. The product of tidal volume (spontaneous and timed) and rate (spontaneous and timed). Moving average over the last 6 breaths.</td>
</tr>
<tr>
<td>$V_T$</td>
<td>Estimated exhaled tidal volume. Moving average over the last 6 breaths. It is body temperature pressure saturated (BTPS) compensated.</td>
</tr>
<tr>
<td><strong>Waveform window</strong></td>
<td></td>
</tr>
<tr>
<td>$P$ waveform</td>
<td>Airway pressure. Where applicable, dotted lines represent target IPAP and EPAP.</td>
</tr>
<tr>
<td>$V$ waveform</td>
<td>Estimated patient flow. The total delivered flow minus the leak flow ($TotLeak$), where $TotLeak$ includes known (intentional) leakage through the exhalation port plus any unintentional leakage in the circuit or at the mask/patient interface.</td>
</tr>
<tr>
<td>V waveform</td>
<td>Estimated patient volume. In AVAPS mode, the dotted line represents target volume.</td>
</tr>
</tbody>
</table>

Scaling the waveform axes

Scale the vertical and horizontal waveform axes with the scale buttons.

The vertical scale button autoscales the Y axes to best fit the current data.

The horizontal (time adjust) button rescales the X axis to show 3, 6, 12, or 24 seconds.
Freezing and unfreezing waveforms

Freeze waveforms for extended viewing by selecting the pause button to the left of the waveform window. The cursor makes one complete sweep across the waveform and then displays the pause in progress symbol. The graphic display is then frozen, and the cursor is visible in the middle of the display (Figure 8-2). Reposition the cursor by touching the waveform screen or with the navigation ring (legacy versions). Data values at cursor location for pressure, flow, and volume are displayed in the white boxes.

Unfreeze the waveforms with the resume button.

![Waveform window with frozen screen](image)

**Figure 8-2: Waveform window with frozen screen**
Patient monitoring
Chapter 9. Alarms, messages, and troubleshooting

Alarms and messages on the ventilator alert you to situations that require your attention. The ventilator can also actuate remote alarms. Figure 9-1 on page 2 shows the visual alarm characteristics. Table 9-3 on page 9-7 summarizes the different types of alarm and tells you how to respond to each.

Responding to alarms

WARNING: If AC power fails and the backup battery is not installed or is depleted, an audible and visual alarm annunciates for at least 2 minutes. Immediately discontinue ventilator use and secure an alternative means of ventilation. As in most ventilators with passive exhalation ports, when power is lost, sufficient air is not provided through the circuit and exhaled air may be rebreathed.

NOTE: If an alarm persists for no apparent reason, discontinue ventilator use and contact Philips.

Respond to an alarm as follows:

1. Approach the patient immediately. Secure sufficient and effective ventilation for the patient. You may silence the alarm if possible.

2. Correct the alarm condition, referring to the alarm messages in Table 9-3.

You can modify alarm settings at any time through the Alarm Settings tab.
Alarms, messages, and troubleshooting

Figure 9-1: Visual alarm indications
## Alarms, messages, and troubleshooting

### Table 9-1: Alarm summary

<table>
<thead>
<tr>
<th>Status</th>
<th>Alarm LED on front panel</th>
<th>Alarm status bar</th>
<th>Alarm message in Alarms list</th>
<th>Audio*</th>
<th>Action required</th>
<th>Remote alarm</th>
</tr>
</thead>
<tbody>
<tr>
<td>No alarms</td>
<td>Off</td>
<td></td>
<td>None</td>
<td>Off</td>
<td>None</td>
<td>Off</td>
</tr>
<tr>
<td>Autoreset alarm</td>
<td>Off</td>
<td>Red (high-priority) or yellow (low-priority)</td>
<td>Background color same as that of active alarm. Message with strike-out text. Alarm icon.</td>
<td></td>
<td>Important information or instructions.</td>
<td></td>
</tr>
<tr>
<td>Informational message</td>
<td>Off</td>
<td>Blue</td>
<td>Blue background color. Informational icon.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low-priority alarm</td>
<td>Off</td>
<td>Yellow</td>
<td>Yellow background color. Alarm icon.</td>
<td>Intermittent tone at an interval of approximately 20 seconds</td>
<td>Respond promptly. Troubleshoot as per Table 9-3.</td>
<td></td>
</tr>
<tr>
<td>High-priority alarm</td>
<td>Flashes</td>
<td>Alternates black and red</td>
<td>Red background color. Alarm icon.</td>
<td>Repeating sequence of 5 tones</td>
<td>Respond immediately to ensure patient safety. Troubleshoot as per Table 9-3.</td>
<td>On</td>
</tr>
<tr>
<td>High-priority alarm – Check Vent</td>
<td>Flashes</td>
<td>Alternates black and red</td>
<td>Red background color. Alarm icon.</td>
<td>Repeating sequence of 5 tones</td>
<td>Respond immediately to ensure patient safety. Do not use equipment that is malfunctioning or that indicates a potential problem until the problem is corrected. Troubleshoot as per Table 9-4.</td>
<td></td>
</tr>
<tr>
<td>High-priority alarm – Vent Inoperative</td>
<td>On continuously</td>
<td>Vent Inoperative screen, including code (Figure 9-2)</td>
<td>Primary alarm (Repeating sequence of 5 tones) or backup alarm (alternating tone for a minimum of 2 minutes)</td>
<td>Continued safe ventilator operation may be in jeopardy. O2 flow and blower operation are disabled. Immediately secure alternative ventilation for the patient. Troubleshoot as per Table 9-5.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Loss of power</td>
<td>Off</td>
<td>Blank</td>
<td>Blank</td>
<td></td>
<td>Immediately secure alternative ventilation for the patient.</td>
<td></td>
</tr>
</tbody>
</table>

* The volume of the primary alarm is the same for low- and high-priority alarms.
Alarms, messages, and troubleshooting

Setting alarm loudness

You can set the alarm loudness from the Menu window (see “Loudness” on page 6-17).
Alarms, messages, and troubleshooting

Silencing alarms

Silence an alarm for 2 minutes by selecting the Alarm Silence button.

The button icon is replaced by this one. A timer shows time remaining in the 2-minute alarm silence period.

Select Alarm Silence again at any time to reset the counter to 2:00 minutes. During patient maneuvers, you can pre-silence audible alarms as desired.

Some alarms cannot be silenced; these are listed in Table 9-3. When a non-silenceable alarm is annunciated, the following is shown.

Resetting alarms

Most alarms reset themselves (autoreset) when the alarm triggering condition is removed, but you must manually reset others. Table 9-3 specifies whether an alarm is autoreset.

Manually resetting alarms

Manually reset an alarm by selecting Alarm Reset.

When an alarm is manually reset, the message is cleared from the Alarms list, any other alarm indications are removed, and the alarm silence is terminated.

If the alarm cannot be manually reset, you see the following:

Clearing autoreset alarms from the Alarms list

Autoreset alarms are shown with text crossed out in the Alarms list.

Clear the message from the Alarms list by selecting Alarm Reset.
Alarms, messages, and troubleshooting

Hiding/displaying alarm messages
To hide an alarm or informational message in the Alarms or Messages list, touch the flashing alarm indicator button or informational message button when up arrows are present. To display messages, touch the flashing alarm indicator or Informational Message button when down arrows are present. Both active and autoreset alarms and informational messages are displayed and hidden.

Symptom-based troubleshooting
To troubleshoot a problem with the ventilator, see Table 9-2. If an alarm message is displayed, see also Table 9-3.

Table 9-2: Symptom-based troubleshooting

<table>
<thead>
<tr>
<th>Symptom</th>
<th>Recommended Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ventilation stops, but the ventilator has power. The ventilator may be in standby or in the Vent Inoperative state. When the ventilator is in the Vent Inoperative state, continued safe ventilator operation may be in jeopardy. O₂ flow and blower operation are disabled.</td>
<td>Check the patient. If the ventilator is in standby, reconnect the patient, and ventilation should resume. If the Alarm LED on the front panel is continuously lit, and/or the Vent Inoperative screen is displayed, provide alternative ventilation. Have the ventilator serviced. For details about the specific Vent Inoperative alarm, see Table 9-5 on page 9-16.</td>
</tr>
<tr>
<td>The touchscreen is unresponsive or does not respond properly. The ventilator continues to function at the selected settings. Patient settings and patient data continue to be visible and accurate, and alarms continue to be annunciated.</td>
<td>Check the patient. If the patient requires a setting change, provide alternative ventilation. Have the ventilator serviced. NOTE: If you cannot shut off the ventilator when the touchscreen is unresponsive, press the ON/Shutdown key, then the Accept (✓) button on the top-right front of the ventilator.</td>
</tr>
<tr>
<td>The ventilator does not switch from battery to AC power.</td>
<td>Check the patient. Make sure the ventilator is connected to AC power. (If the ventilator is connected to AC power and the ventilator is functioning correctly, the ON/Shutdown LED should be on.) If problem persists, provide alternative ventilation. Have the ventilator serviced.</td>
</tr>
<tr>
<td>The ventilator does not switch from AC power to battery. A back-up alarm should annunciate.</td>
<td>Check the patient. If problem persists, provide alternative ventilation. Have the ventilator serviced.</td>
</tr>
<tr>
<td>The estimated exhaled tidal volume reading (Vₜ) is inaccurate.</td>
<td>Check the patient. Check for large leaks. Make sure ventilator and alarm settings are appropriate. Make sure an approved patient circuit is in use. Adjust the mask to ensure proper fit and adequate leak compensation. If the problem persists, provide alternative ventilation. Have the ventilator serviced.</td>
</tr>
</tbody>
</table>
Alarms and other messages

Table 9-3 is a list of alarms and other messages displayed by the ventilator, along with descriptions, suggested corrective actions, and other information. The ID (identifier) listed with the priority type is the priority number of the alarm. This priority number determines the order of alarm message display. Unless otherwise indicated, alarms listed as autoresettable are reset when the alarm condition is removed.

<table>
<thead>
<tr>
<th>Message</th>
<th>Description</th>
<th>Corrective Action</th>
<th>Priority type (ID)</th>
<th>Manually resettable</th>
<th>Autoresettable</th>
<th>Silence able</th>
</tr>
</thead>
<tbody>
<tr>
<td>AVAPS: Target VT Exceeded. Min Pressure Too High</td>
<td>AVAPS target pressure is less than Min P setting. The ventilator limits its applied pressure to Min P.</td>
<td>Check the patient. Confirm pressure settings are compatible with target. Evaluate pressure and volume settings.</td>
<td>Information (63)</td>
<td>No</td>
<td>Yes</td>
<td>N/A</td>
</tr>
<tr>
<td>AVAPS: Target VT Not Achieved. Insufficient Max Pressure</td>
<td>AVAPS target pressure exceeds Max P setting. The ventilator limits applied pressure to Max P.</td>
<td>Check the patient. Confirm pressure settings are compatible with target. Evaluate pressure and volume settings.</td>
<td>Information (62)</td>
<td>No</td>
<td>Yes</td>
<td>N/A</td>
</tr>
<tr>
<td>Bacteria filter must be installed onto gas outlet</td>
<td>An inspiratory bacteria filter must be installed on the patient gas outlet port.</td>
<td>Confirm that a bacteria filter is installed. Install a bacteria filter if one is not present.</td>
<td>Information (67)</td>
<td>Yes</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Cannot Reach Target Flow</td>
<td>Displays when HFT (high flow therapy) is active. Indicates that flow target is not achieved.</td>
<td>Check the patient. Check that the high flow nasal cannula size is appropriate for the flow setting. Check that an occlusive interface is NOT in use (a cannula fully sealed within the nares, an NIV mask or direct connection to an ETT/trach). Check for an occlusion, kink or liquid in the patient circuit.</td>
<td>Low (53)</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Check Vent: description of failure</td>
<td>See Table 9-4 on page 9-12</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>High Inspiratory Pressure</td>
<td>Measured inspiratory pressure is greater than the HIP setting, and the ventilator cycles into exhalation. Autoresets after a complete inspiration without the alarm condition.</td>
<td>Check the patient. Confirm ventilator and alarm settings are appropriate. If problem persists, provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (45)</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>High O₂ Supply Pressure</td>
<td>O₂ inlet pressure is greater than 92 psig, so O₂ enrichment ends. Autoresets when O₂ supply pressure falls below 87 psig.</td>
<td>Check the patient. If problem persists, provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (49)</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
</tr>
</tbody>
</table>
## Alarms, messages, and troubleshooting

<table>
<thead>
<tr>
<th>Message</th>
<th>Description</th>
<th>Corrective Action</th>
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</tr>
</thead>
<tbody>
<tr>
<td>High Rate</td>
<td>Measured respiratory rate is greater than the Hi Rate setting. Escalates to a high-priority alarm if the alarm condition persists for more than 60 sec.</td>
<td>Check the patient. Confirm ventilator and alarm settings are appropriate. If problem persists, provide alternative ventilation. Have the ventilator serviced.</td>
<td>Low/High (59)</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>High Tidal Volume</td>
<td>Measured estimated tidal volume is greater than the Hi VT setting. Escalates to a high-priority alarm if the alarm condition persists for more than 60 sec.</td>
<td>Check the patient. Check for large leaks. Confirm ventilator and alarm settings are appropriate. If problem persists, provide alternative ventilation. Have the ventilator serviced.</td>
<td>Low/High (58)</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Low Inspiratory Pressure</td>
<td>Measured inspiratory pressure is less than the LIP setting.</td>
<td>Check the patient. Confirm ventilator and alarm settings are appropriate. If problem persists, provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (46)</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Low Internal Battery</td>
<td>Battery can provide operating power for only an additional 15 minutes under nominal conditions. Autoresets when ventilator is connected to AC power.</td>
<td>Connect ventilator to AC power. Provide alternative ventilation.</td>
<td>High (43)</td>
<td>No</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Low Leak–CO₂ Rebreathing Risk</td>
<td>Estimated volume of exhaled gas returned to the patient is high.</td>
<td>Check the patient, as possibility of CO₂ rebreathing could pose a potential problem. Check the exhalation port for occlusions. Check for appropriate patient interface and exhalation port settings. If the approved exhalation port is unobstructed, mask and port settings are appropriate, and problem persists, increase the ventilator baseline flow by adding leak or increasing EPAP, if possible. Switch to an approved nebulizer.</td>
<td>High (42)</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
</tbody>
</table>
### Table 9-3: Alarm and other messages: summary and troubleshooting (continued)

<table>
<thead>
<tr>
<th>Message</th>
<th>Description</th>
<th>Corrective Action</th>
<th>Priority type (ID)</th>
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<th>Silence able</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low Minute Ventilation</td>
<td>Estimated minute ventilation is less than the Lo $V_E$ setting. Escalates to a high-priority alarm if the alarm condition persists for more than 60 sec.</td>
<td>Check the patient. Confirm ventilator and alarm settings are appropriate. If problem persists, provide alternative ventilation. Have the ventilator serviced.</td>
<td>Low (56)</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Low $O_2$ Supply Pressure</td>
<td>$O_2$ inlet pressure is less than 30 psig and delivered $O_2$ is at least 5% lower than $O_2$ setting. The ventilator continues to deliver as much $O_2$ support as possible, but it ends $O_2$ support when the $O_2$ inlet pressure drops to less than 18 psig. Autoresets when $O_2$ inlet pressure exceeds 23 psig.</td>
<td>Check the patient. Attach to oxygen source with sufficient pressure. If problem persists, provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (48)</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
</tr>
</tbody>
</table>
| Low Rate                       | A low-priority alarm if the measured respiratory rate is less than the Lo Rate setting, escalating to a high-priority alarm in 60 sec. A high-priority alarm from the start if:  
  - The Lo Rate setting is $\leq$ 4 BPM and there are no breaths for $>60$/Lo Rate setting.  
  - The Lo Rate setting is $> 4$ BPM and there are no breaths for $>15$ sec. | Check the patient. Confirm ventilator and alarm settings are appropriate. If problem persists, provide alternative ventilation. Have the ventilator serviced. | Low/High (55)      | Yes                | Yes            | Yes          |
| Low Tidal Volume               | Estimated tidal volume is less than the Lo $V_T$ setting. Escalates to a high-priority alarm if the alarm condition persists for more than 60 sec. | Check the patient. Confirm ventilator and alarm settings are appropriate. If problem persists, provide alternative ventilation. Have the ventilator serviced. | Low (57)           | Yes                | Yes            | Yes          |
| Mask: x, Exh Port: y Use Menu to change | Displays when ventilator is turned on. Displays selected mask type and exhalation port. | Select mask and port from Menu tab. Message is removed when user confirms selections, or after 5 minutes. | Information (64)   | No                 | Yes            | N/A          |
### Alarms, messages, and troubleshooting

Table 9-3: Alarm and other messages: summary and troubleshooting (continued)

<table>
<thead>
<tr>
<th>Message</th>
<th>Description</th>
<th>Corrective Action</th>
<th>Priority type (ID)</th>
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<th>Autoresettable</th>
<th>Silenceable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oxygen Not Available</td>
<td>Oxygen supply pressure out of range, oxygen device failed, air flow sensor and/or oxygen flow sensor calibration failed, or oxygen inlet pressure sensor calibration failed. The ventilator discontinues oxygen support.</td>
<td>Check the patient. Check if high/low O₂ source is the problem and correct. If problem persists, provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (47)</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Patient alarms are disabled during HFT</td>
<td>Displays when HFT (high flow therapy) is active. Patient alarms are not available in this therapy.</td>
<td>Manually reset to confirm and clear the audible alarm.</td>
<td>Low/Information (54)</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Patient Circuit Occluded</td>
<td>Proximal pressure and patient flow are low. Patient circuit occluded.</td>
<td>Check the patient. Check the patient circuit for bulk liquid, crimps, or blocked filter. Confirm ventilator and alarm settings are appropriate. If problem persists, provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (40)</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Patient Circuit Occluded</td>
<td>Displays when HFT (high flow therapy) is active. Gas flow to the patient is obstructed.</td>
<td>Check the patient. Check that an occlusive interface is NOT in use (a cannula fully sealed within the nares, an NIV mask or direct connect to an ETT/trach). Check for an occlusion, kink or liquid in the patient circuit. If problem persists, provide alternative ventilation.</td>
<td>High (52)</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Patient Disconnect</td>
<td>Patient is no longer connected to the ventilator, either through circuit or mask, or the patient circuit is disconnected from the ventilator and the patient is no longer receiving ventilatory support. Ventilation continues. NOTE: The Patient Disconnect alarm is triggered when 11 seconds have elapsed.</td>
<td>Check the patient. Reconnect patient circuit. Confirm ventilator and alarm settings are appropriate. If problem persists, provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (39)</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
</tbody>
</table>
# Alarms, messages, and troubleshooting

## Table 9-3: Alarm and other messages: summary and troubleshooting (continued)

<table>
<thead>
<tr>
<th>Message</th>
<th>Description</th>
<th>Corrective Action</th>
<th>Priority type (ID)</th>
<th>Manually resettable</th>
<th>Autoresettable</th>
<th>Silence able</th>
</tr>
</thead>
<tbody>
<tr>
<td>Power has been re-</td>
<td>Power is restored following loss of power. The ventilator restarts and</td>
<td>Check the patient. Confirm ventilator and alarm settings are appropriate.</td>
<td>Information (66)</td>
<td>Yes</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>stored</td>
<td>continues ventilation in the mode set before power was lost.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PPV Maximum Pressure</td>
<td>Computed target pressure is greater than the PPV maximum pressure alarm</td>
<td>Check the patient. Confirm ventilator and alarm settings are appropriate.</td>
<td>Information/High</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td>limit. Possible causes are excessive patient inspiratory effort; a significant</td>
<td>Check for circuit or mask leaks. If problem persists, provide alternative ventilation. Have the ventilator serviced.</td>
<td>(51)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>change in the leak around the patient interface; or high PPV %, Max E, or Max</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>R setting. Target pressure is limited.</td>
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<td></td>
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<td></td>
</tr>
<tr>
<td></td>
<td>At first, an information message. If condition persists for three consecutive</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>PPV inspirations, this escalates to a high-priority alarm.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PPV Maximum Volume</td>
<td>Estimated delivered patient tidal volume is greater than the PPV maximum</td>
<td>Check the patient. Confirm ventilator and alarm settings are appropriate.</td>
<td>Information/High</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td>volume alarm limit. Possible causes are excessive patient inspiratory effort;</td>
<td>Check for circuit or mask leaks. If problem persists, provide alternative ventilation. Have the ventilator serviced.</td>
<td>(50)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>a significant change in the leak around the patient interface; or high PPV</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>%, Max E, or Max R setting. Ventilator cycles to exhalation.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>At first, an information message. If condition persists for three consecutive</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>PPV inspirations, this escalates to a high-priority alarm.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pressure Regulation</td>
<td>Pressures exceed ventilator-defined thresholds for IPAP, EPAP, or both.</td>
<td>Check the patient. Confirm ventilator and alarm settings are appropriate.</td>
<td>High (44)</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>High</td>
<td>Ventilation continues. Autoresets when alarm condition removed; otherwise, transitions to the ventilator in-operative state if pressure continues to rise.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Alarms, messages, and troubleshooting

Table 9-3: Alarm and other messages: summary and troubleshooting (continued)

<table>
<thead>
<tr>
<th>Message</th>
<th>Description</th>
<th>Corrective Action</th>
<th>Priority type (ID)</th>
<th>Manually resettable</th>
<th>Autoresettable</th>
<th>Silenceable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Proximal Pressure Line Disconnect</td>
<td>Proximal pressure low for a few seconds. Proximal pressure line is disconnected. Air flow to the patient continues.</td>
<td>Check the patient. Reconnect proximal pressure line. Confirm ventilator and alarm settings are appropriate. If problem persists, provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (41)</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Running on Internal Battery</td>
<td>System is powered by the internal battery. Autoresets when ventilator is connected to AC power.</td>
<td>Connect ventilator to AC power.</td>
<td>Low (60)</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Trigger:+x, E-Cycle: +x Use Menu to change</td>
<td>Auto-Trak+ is active and using the displayed settings. This message is displayed for 5 min after start-up.</td>
<td>Confirm that Auto-Trak+ settings are appropriate.</td>
<td>Information (65)</td>
<td>Yes</td>
<td>Yes</td>
<td>N/A</td>
</tr>
<tr>
<td>Using Default Settings</td>
<td>Displayed after power on if setting values are corrupted or not set, or if default values were restored by the user.</td>
<td>Check the patient. Check and adjust settings as required.</td>
<td>Information (61)</td>
<td>Yes</td>
<td>Yes</td>
<td>N/A</td>
</tr>
<tr>
<td>Vent Inoperative x description of failure</td>
<td>See Table 9-5 on page 9-16</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 9-4: Check Vent alarm messages: summary and troubleshooting

<table>
<thead>
<tr>
<th>Message</th>
<th>Description</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Check Vent: 1.8 V Supply Failed</td>
<td>Technical failure</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (21)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: 3.3 V Supply Failed</td>
<td>Technical failure</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (22)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: 5 V Supply Failed</td>
<td>Technical failure</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (23)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: 12 V Supply Failed</td>
<td>Technical failure</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (24)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: 24 V Supply Failed</td>
<td>Technical failure</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (25)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>
### Alarms, messages, and troubleshooting

Table 9-4: Check Vent alarm messages: summary and troubleshooting (continued)

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<tr>
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<th>Silenceable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Check Vent: 35 V Supply Failed</td>
<td>Technical failure</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (26)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Air Flow Sensor Calibration Data Error</td>
<td>Flow-related patient data is disabled. O₂ concentration switches to 21% (ventilates with air only). Default volume used in AVAPS mode. Standby disabled. Volume, leak, disconnect, and occlusion alarms compromised.</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (13)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Alarm LED Failed</td>
<td>Technical failure.</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (6)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Aux Supply Failed</td>
<td>Backup alarm problem</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (20)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Backup Alarm Failed</td>
<td>Backup alarm problem</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (5)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Barometer Calibration Data Error</td>
<td>Default barometric pressure of 686.0 mmHg (approximately 900 m/2953 ft above sea level) used in calculations</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (18)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Barometer Sensor Range Error</td>
<td>Default barometric pressure of 686.0 mmHg (approximately 900 m/2953 ft above sea level) used in calculations</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (19)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Battery Failed</td>
<td>Battery problem</td>
<td>Check the patient. Connect the ventilator to AC. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (35)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Battery Temperature High</td>
<td>Battery problem</td>
<td>Check the patient. Connect the ventilator to AC. Check for causes of overheating, such as high room temperature, blocked vents, clogged air inlet filter, or nonfunctional fan. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (34)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>
### Alarms, messages, and troubleshooting

Table 9-4: Check Vent alarm messages: summary and troubleshooting (continued)

<table>
<thead>
<tr>
<th>Message</th>
<th>Description</th>
<th>Corrective Action</th>
<th>Priority type (ID)</th>
<th>Manually resettable</th>
<th>Autoresettable</th>
<th>Silenceable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Check Vent: Blower Stalled</td>
<td>Technical failure</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (2)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Blower Temperature High</td>
<td>Technical failure</td>
<td>Check the patient. Check for causes of overheating, such as high room temperature, blocked vents, clogged air inlet filter, or nonfunctional fan. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (33)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Cooling Fan Speed Error</td>
<td>Overheating of ventilator possible</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (36)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: CPU PCBA ADC Failed</td>
<td>Technical failure</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (29)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Data Acquisition PCBA ADC Failed</td>
<td>Technical failure</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (27)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Flash File System Error</td>
<td>Technical failure</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (37)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Internal Temperature High CPU</td>
<td>Technical failure</td>
<td>Check the patient. Check for causes of overheating, such as high room temperature, blocked vents, clogged air inlet filter, or nonfunctional fan. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (30)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Internal Temperature High Daq</td>
<td>Technical failure</td>
<td>Check the patient. Check for causes of overheating, such as high room temperature, blocked vents, clogged air inlet filter, or nonfunctional fan. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (31)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Internal Temperature High Mtr</td>
<td>Technical failure</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (32)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Machine Pressure Sensor Auto-Zero Failed</td>
<td>Proximal pressure is not measured. Pressure-related alarms are compromised.</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (9)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>
### Table 9-4: Check Vent alarm messages: summary and troubleshooting (continued)

<table>
<thead>
<tr>
<th>Message</th>
<th>Description</th>
<th>Corrective Action</th>
<th>Priority type (ID)</th>
<th>Manually resettable</th>
<th>Autoresettable</th>
<th>Silenceable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Check Vent: Machine Pressure Sensor Calibration Data Error</td>
<td>Proximal pressure is not measured. Pressure-related alarms are compromised.</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (7)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Machine Pressure Sensor Range Error</td>
<td>Proximal pressure is not measured. Pressure-related alarms are compromised.</td>
<td></td>
<td>High (11)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Motor Control PCBA ADC Failed</td>
<td>Technical failure</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (28)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: O₂ Flow Sensor Calibration Data Error</td>
<td>Continues to ventilate with air only</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (14)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: O₂ Pressure Sensor Calibration Data Error</td>
<td>Continues to ventilate with air only</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (15)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: O₂ Supply Pressure Sensor Range Error</td>
<td>Continues to ventilate with air only</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (17)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: OVP Circuit Failed</td>
<td>Technical failure</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (38)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Oxygen Device Failed</td>
<td>Continues to ventilate with air only</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (16)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Primary Alarm Failed</td>
<td>Technical failure</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (4)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Program CRC Test Failed</td>
<td>Technical failure</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (1)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Proximal Pressure Sensor Auto-Zero Failed</td>
<td>Proximal pressure is not measured. Pressure-related alarms are compromised.</td>
<td></td>
<td>High (10)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Proximal Pressure Sensor Calibration Data Error</td>
<td>Proximal pressure is not measured. Pressure-related alarms are compromised.</td>
<td></td>
<td>High (8)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>
## Alarms, messages, and troubleshooting

### Table 9-4: Check Vent alarm messages: summary and troubleshooting (continued)

<table>
<thead>
<tr>
<th>Message</th>
<th>Description</th>
<th>Corrective Action</th>
<th>Priority type (ID)</th>
<th>Manually resettable</th>
<th>Autoresettable</th>
<th>Silenceable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Check Vent: Proximal Pressure Sensor Range Error</td>
<td>Proximal pressure is not measured. Pressure-related alarms are compromised. Also displays when HFT (high flow therapy) is active and proximal pressure is out of range.</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced. When using a high flow nasal cannula with FEP Connect, the proximal pressure line should be disconnected from the ventilator port during HFT. If this message is seen when switching the patient from ventilation to HFT, check to make sure the proximal pressure line is disconnected.</td>
<td>High (12)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Check Vent: Ventilator Restarted</td>
<td>Technical failure</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (3)</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>

### Table 9-5: Vent Inoperative alarm messages: summary and troubleshooting

<table>
<thead>
<tr>
<th>Message</th>
<th>Description</th>
<th>Corrective Action</th>
<th>Priority type (ID)</th>
<th>Manually resettable</th>
<th>Autoresettable</th>
<th>Silenceable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vent Inoperative 1000 3.3 V Supply Failed</td>
<td>Technical failure. The ventilator is in the ventilator inoperative state.</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (2)</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Vent Inoperative 1001 12 V Supply Failed</td>
<td>Technical failure. The ventilator is in the ventilator inoperative state.</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (3)</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Vent Inoperative 1002 Blower Temperature Too High</td>
<td>Technical failure. The ventilator is in the ventilator inoperative state.</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (4)</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Vent Inoperative 1003 Internal Temperature High</td>
<td>Technical failure of the CPU PCBA. The ventilator is in the ventilator inoperative state.</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (5)</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Vent Inoperative 1004 Internal Temperature High</td>
<td>Technical failure of the DAQ PCBA. The ventilator is in the ventilator inoperative state.</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (6)</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>
## Alarms, messages, and troubleshooting

### Table 9-5: Vent Inoperative alarm messages: summary and troubleshooting (continued)

<table>
<thead>
<tr>
<th>Message</th>
<th>Description</th>
<th>Corrective Action</th>
<th>Priority type (ID)</th>
<th>Manually resettable</th>
<th>Autoresettable</th>
<th>Silenceable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vent Inoperative 1005 Internal Temperature High</td>
<td>Technical failure of the motor PCBA. The ventilator is in the ventilator inoperative state.</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (7)</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Vent Inoperative 1006 Data Acquisition PCBA ADC Failed</td>
<td>Technical failure. The ventilator is in the ventilator inoperative state.</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (8)</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Vent Inoperative 1007 Machine and Proximal Pressure Sensors Failed</td>
<td>Technical failure. The ventilator is in the ventilator inoperative state.</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (9)</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Vent Inoperative 1008 Machine and Proximal Pressure Sensors Failed</td>
<td>Technical failure. The ventilator is in the ventilator inoperative state.</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (10)</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Vent Inoperative 1009 Pressure Regulation High</td>
<td>Technical failure. The ventilator is in the ventilator inoperative state.</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (11)</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Vent Inoperative 100A Data Acquisition PCBA ADC Reference Failed</td>
<td>Technical failure. The ventilator is in the ventilator inoperative state.</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (12)</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Vent Inoperative 100B Watchdog Test Failed</td>
<td>Technical failure. The ventilator is in the ventilator inoperative state.</td>
<td>Check the patient. Provide alternative ventilation. Have the ventilator serviced.</td>
<td>High (13)</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>
Alarms, messages, and troubleshooting
Chapter 10. Care and maintenance

WARNING: To reduce the risk of electric shock, power down the ventilator and disconnect it from AC power before cleaning, disinfecting, or servicing it.

WARNING: This product consists of devices that may contain mercury, which must be recycled or disposed of in accordance with local, state, or federal laws. (Within this system, the backlight lamps in the monitor display contain mercury.)

WARNING: Before performing maintenance procedures, disconnect the ventilator from the patient, shut it down, and disconnect it from AC power. All operator maintenance must be performed with the patient off the ventilator. Failure to do so can result in electric shock to the patient and operator.

NOTE: It is the user’s responsibility to comply with the information provided in this chapter.

NOTE: Cleaning and disinfection are most effective if soiling is not allowed to dry on a medical device.¹

NOTE: Disinfection is most effective on medical devices that were previously cleaned.¹

NOTE: For all V60/V60 Plus hardware accessories recommended by Philips, follow the cleaning and disinfection guidelines in this chapter. For multi-patient interface and circuit accessories, consult the product instructions for use. For single patient use accessories, no cleaning and disinfection is needed.

To ensure the safety and reliability of your ventilator, follow these maintenance procedures along with your own institutional policies for cleaning, disinfecting, and maintaining equipment. All the procedures in this manual are intended to be performed by the operator. For further maintenance, contact your service representative.

¹ Reprocessing Medical Devices in Health Care Settings: Validation Methods and Labeling Guidance for Industry and Food and Drug Administration Staff, 2015. Food and Drug Administration (FDA)
Care and maintenance

Exterior and touchscreen cleaning

CAUTION: To prevent possible damage to the ventilator, use only those cleaning and disinfecting agents listed in this manual.

CAUTION: To prevent possible damage to the ventilator, do not drip or spray any liquids directly onto any surface including the front panel, touchscreen, navigation ring (legacy versions), and Accept button.

CAUTION: Never clean or disinfect the touchscreen with an abrasive brush or device, since this will cause irreparable damage.

CAUTION: Do not attempt to sterilize or autoclave the ventilator.

NOTE: Use of unapproved cleaning and disinfecting agents may cause damage to the enclosure, touchscreen, or parts of the ventilator.

Approved cleaning agents

The following cleaning agent is acceptable for use on the touchscreen and exterior surfaces of the ventilator:

- Medivators Intercept Detergent, per manufacturer's recommendation at 1/3 oz (10 mL) per gallon of warm tap water.

Cleaning instructions

1. Apply cleaning agent to a soft lint-free cloth or use a disposable wipe. The cloth or wipe should be saturated but not dripping.

2. Wipe cleaning agent over the entire exterior surface and touchscreen of the ventilator.

3. Continue wiping until all visible contaminants and soiling are removed.

4. Rinse with a clean, water-dampened cloth and allow to dry completely before reuse.
Exterior and touchscreen disinfection

CAUTION: To prevent possible damage to the ventilator, use only those cleaning and disinfecting agents listed in this manual.

CAUTION: To prevent possible damage to the ventilator, do not drip or spray any liquids directly onto any surface including the front panel, touchscreen, navigation ring (legacy versions), and Accept button.

CAUTION: Never clean or disinfect the touchscreen with an abrasive brush or device, since this will cause irreparable damage.

CAUTION: Do not attempt to sterilize or autoclave the ventilator.

NOTE: Use of unapproved cleaning and disinfecting agents may cause damage to the enclosure, touchscreen, or parts of the ventilator.

Approved disinfecting agents

The following disinfecting agents are acceptable for use on the touchscreen and exterior surfaces of the ventilator:

Table 10-1: Exterior disinfection

<table>
<thead>
<tr>
<th>Disinfectant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Solution of 1 part 5% sodium hypochlorite (bleach) diluted in 9 parts deionized water.</td>
</tr>
<tr>
<td>3% hydrogen peroxide</td>
</tr>
</tbody>
</table>

Disinfection instructions

1. Ensure that cleaning was done per the procedure in “Exterior and touchscreen cleaning” on page 10-2.

2. Apply disinfecting agent to a soft lint-free cloth or use a disposable wipe. The cloth or wipe should be saturated but not dripping.

3. Wipe disinfecting agent over the entire exterior surface of the ventilator.

4. Allow disinfectant to remain on the surface for the following contact times:
   - bleach solution — 2 minutes
   - hydrogen peroxide — 15 minutes

5. Rinse with a clean cloth dampened with water and allow to dry completely before reuse.
Care and maintenance

**Bacteria filter, patient circuit, and other accessories**

Follow the manufacturer’s instructions that accompany the accessory.

---

**WARNING:** To prevent patient or ventilator contamination, inspect and replace the main flow bacteria filter between patients and at regular intervals (or as stated by the manufacturer).

**WARNING:** To prevent possible patient injury, inspect and verify the proper operation of the exhalation port regularly during use.

**CAUTION:** Because some environments cause a quicker collection of lint and dust than others, inspect the filters more often when needed. The air inlet filter should be replaced; the cooling fan filter should be cleaned.

**CAUTION:** To ensure proper system performance, use a Respironics-approved air inlet filter.
Preventive maintenance

**WARNING:** Only authorized service personnel should replace parts within the ventilator or perform other service activities. Unauthorized personnel without proper training are at risk of electric shock.

**WARNING:** Turn off the ventilator and disconnect it from the AC mains outlet before you perform decontamination or maintenance procedures. Failure to do so may result in electric shock.

The expected service life of your V60/V60 Plus ventilator is 10 years. Perform preventive maintenance on the ventilator according to the schedule in Table 10-2. You can view the hours of ventilator operation in the Vent Info window (“Vent Info (ventilator information)” on page 6-18). The following subsections provide details for some of these preventive maintenance procedures.

<table>
<thead>
<tr>
<th>Frequency</th>
<th>Component</th>
<th>Maintenance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Between patients and per institutional guidelines</td>
<td>Patient circuit</td>
<td>Per manufacturer recommendations.</td>
</tr>
<tr>
<td></td>
<td>Main flow bacteria filter</td>
<td>Replace per institutional guidelines.</td>
</tr>
<tr>
<td>Every month</td>
<td>Cooling fan filter</td>
<td>Inspect for occlusions, dust, lint, etc. If discolored or dirty, remove and wash or rinse thoroughly, and let dry completely before reinstalling.</td>
</tr>
<tr>
<td></td>
<td>Air inlet filter</td>
<td>Inspect and replace if needed</td>
</tr>
<tr>
<td>Every year</td>
<td>Backup battery</td>
<td>Inspect, test, and replace if needed*</td>
</tr>
<tr>
<td></td>
<td>Ventilator</td>
<td>Preventive maintenance*</td>
</tr>
<tr>
<td>As required</td>
<td>Backup battery</td>
<td>A new backup battery should be installed and charged within one year of the date of manufacture identified on the battery and on the shipping box.</td>
</tr>
<tr>
<td>Every 5 years</td>
<td>Backup battery</td>
<td>Replace.* Battery replacement is based on the date of manufacture recorded on the battery label. Also viewable in Diagnostic Mode on the system information screen.</td>
</tr>
</tbody>
</table>

* Must be done by authorized service personnel according to the instructions in the service manual.
Re replacing the air inlet filter
Replace the air inlet filter as follows, referring to Figure 10-1.

1. Power down the ventilator and disconnect it from AC power. Remove ventilator from cart, if applicable.

2. Turn the captive D-ring fastener counter-clockwise one-quarter turn and release. Remove the side panel.

3. Remove the inlet filter by pinching it out of the recess in the bracket.

4. Install a new air filter by tucking it into the recessed area. Replace the side panel, and push in and turn the D-ring fastener one-quarter turn until it locks.

Figure 10-1: Replacing the air inlet filter
Cleaning or replacing the cooling fan filter

Clean or replace the cooling fan filter as follows, referring to Figure 10-2:

1. Insert a small, flat blade driver tip between the foam filter and the filter retaining cover (Figure 10-2).
2. Gently pry the filter cover from the back of the ventilator. Do not remove the fan retaining pins.
3. Wash or rinse the filter. Let it dry completely before reinstalling.
4. Replace the filter, then snap the filter cover into place.

Figure 10-2: Replacing the cooling fan filter
Care and maintenance

Removing and replacing the battery
See “Installing the battery” on page A-4.

Disposal

WARNING: This product consists of devices that may contain mercury, which must be recycled or disposed of in accordance with local, state, or federal laws. (Within this system, the backlight lamps in the monitor display contain mercury.)

Dispose of all parts removed from the device according to your institution’s protocol. Follow all local, state, and federal regulations with respect to environmental protection, especially when disposing of the electronic device or parts of it (for example, oxygen cell, batteries).

Dispose of this device in accordance with local collections and recycling regulations. For more information, visit www.philips.com/recycling.

This device contains a lithium ion battery that cannot be recycled. Please contact customer service for safe disposal instructions of this part.

Storage between patient use

Follow the steps below when storing the ventilator between patient use:

1. Ensure that the patient circuit is assembled and installed properly as described in “Installing the patient circuit” on page 5-3.
2. Plug the ventilator into an AC outlet and verify that the power source symbol is displayed.
3. Adjust settings to hospital defaults.
4. Check oxygen cylinder fill status (if applicable).
5. Ensure oxygen cylinders are turned off.
6. Ensure the environmental specifications are met. Refer to Table 11-8: “Environmental specifications” on page 11-6.

CAUTION: Avoid allowing the ventilator battery to become completely discharged. Otherwise, the battery may become over-discharged and require long recharge times of up to 16 hours or more. The over-discharged condition may permanently damage the battery so that it is unable to recharge. To prevent the occurrence of a non-recoverable over-discharged battery, always keep the ventilator connected to an AC outlet while in storage and schedule regular preventive maintenance to ensure battery health.
Service and repairs

For technical service or repair information not included in this chapter, contact Philips.

A V60/V60 Plus service manual is available (order number 989805612651, part number 1049766), and can also be downloaded from www.philips.com/hrcmanuals. The service manual includes removal and installation procedures, parts lists, and testing and troubleshooting information.

Repacking and shipping

CAUTION: To prevent possible damage to the ventilator, always ship it with the original packing material. If the original material is not available, contact Philips to order replacements.

NOTE: Transport of lithium ion batteries is strictly controlled by international regulations and laws. Do not ship the battery either in the ventilator or separately by sea or air.

Remove the battery from the ventilator before shipping the ventilator. See “Installing the battery” on page A-4 for more information. Ship the battery and ventilator separately in appropriate packaging in conformance with federal, state, and local regulations.

If necessary, Philips will send you a battery return kit that contains a prepaid postal label and instructions for returning the battery if it has failed prematurely (within 5 years of its manufacture date). The order number for the return kit is 989805663471 / part number 1146145.
Care and maintenance
Chapter 11. Technical specifications

Control settings  
Table 11-1 lists ventilator control setting ranges, resolutions, and accuracies. Table 11-2 lists the controls active in the different ventilation modes.

Table 11-1: Control setting ranges, resolutions, and accuracies

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Range</th>
<th>Resolution</th>
<th>Performance Accuracy</th>
<th>Factory default</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Mode settings</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Modes</td>
<td>AVAPS, CPAP, S/T, PCV, PPV (optional)</td>
<td>N/A</td>
<td>N/A</td>
<td>S/T</td>
</tr>
<tr>
<td><strong>Control settings</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C-Flex</td>
<td>OFF, 1 to 3</td>
<td>1</td>
<td>N/A</td>
<td>2</td>
</tr>
<tr>
<td>CPAP</td>
<td>4 to 25 cmH\textsubscript{2}O</td>
<td>1 cmH\textsubscript{2}O</td>
<td>± (2 cmH\textsubscript{2}O + 4% of target)</td>
<td>4 cmH\textsubscript{2}O</td>
</tr>
<tr>
<td>EPAP</td>
<td>4 to 25 cmH\textsubscript{2}O</td>
<td>1 cmH\textsubscript{2}O</td>
<td>± (2 cmH\textsubscript{2}O + 4% of target)</td>
<td>4 cmH\textsubscript{2}O</td>
</tr>
<tr>
<td>Flow (High flow therapy, 3.00 software and above, and V60 Plus)</td>
<td>10 to 80 L/min</td>
<td>5 L/min</td>
<td>N/A</td>
<td>35 L/min</td>
</tr>
<tr>
<td>IPAP</td>
<td>4 to 40 cmH\textsubscript{2}O</td>
<td>1 cmH\textsubscript{2}O</td>
<td>± (2 cmH\textsubscript{2}O + 4% of target)</td>
<td>12 cmH\textsubscript{2}O</td>
</tr>
<tr>
<td>i-Time (Inspiratory Time)</td>
<td>0.30 to 3.00 sec</td>
<td>0.05 sec</td>
<td>± 0.03 sec</td>
<td>1.00 sec</td>
</tr>
<tr>
<td>Max E</td>
<td>0 to 100 cmH\textsubscript{2}O/L</td>
<td>1 cmH\textsubscript{2}O/L</td>
<td>N/A</td>
<td>15 cmH\textsubscript{2}O/L</td>
</tr>
<tr>
<td>Max P (PPV Maximum Pressure Limit)</td>
<td>5 to 40 cmH\textsubscript{2}O</td>
<td>1 cmH\textsubscript{2}O</td>
<td>± (2 cmH\textsubscript{2}O + 4% of target)</td>
<td>20 cmH\textsubscript{2}O</td>
</tr>
<tr>
<td>Max P (AVAPS Maximum IPAP Pressure)</td>
<td>6 to 40 cmH\textsubscript{2}O</td>
<td>1 cmH\textsubscript{2}O</td>
<td>± (2 cmH\textsubscript{2}O + 4% of target)</td>
<td>25 cmH\textsubscript{2}O</td>
</tr>
<tr>
<td>Max R</td>
<td>0 to 50 cmH\textsubscript{2}O/L/sec</td>
<td>1 cmH\textsubscript{2}O/L/s</td>
<td>N/A</td>
<td>4 cmH\textsubscript{2}O/L/s</td>
</tr>
<tr>
<td>Max V (PPV Maximum Volume Limit)</td>
<td>200 to 3500 mL</td>
<td>5 mL</td>
<td>± 15%</td>
<td>1000 mL</td>
</tr>
<tr>
<td>Min P (AVAPS Minimum IPAP Pressure)</td>
<td>5 to 30 cmH\textsubscript{2}O</td>
<td>1 cmH\textsubscript{2}O</td>
<td>± (2 cmH\textsubscript{2}O + 4% of target)</td>
<td>10 cmH\textsubscript{2}O</td>
</tr>
<tr>
<td>O\textsubscript{2} (Oxygen)</td>
<td>21 to 100%</td>
<td>1%</td>
<td>± 5%</td>
<td>21%</td>
</tr>
<tr>
<td>PPV %</td>
<td>0 to 100%</td>
<td>1%</td>
<td>N/A</td>
<td>30%</td>
</tr>
</tbody>
</table>
### Technical specifications

#### Table 11-1: Control setting ranges, resolutions, and accuracies (continued)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Range</th>
<th>Resolution</th>
<th>Performance Accuracy</th>
<th>Factory default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ramp Time</td>
<td>OFF, 5 to 45 min</td>
<td>5 min</td>
<td>± 1 sec</td>
<td>OFF</td>
</tr>
<tr>
<td>Rate (Respiratory Rate)</td>
<td>4 to 60 BPM</td>
<td>1 BPM</td>
<td>± 1 BPM</td>
<td>4 BPM</td>
</tr>
<tr>
<td>Rise (Rise Time)</td>
<td>1 to 5</td>
<td>1</td>
<td>N/A</td>
<td>3</td>
</tr>
<tr>
<td>VT (AVAPS Target Tidal Volume)</td>
<td>200 to 2000 mL BTPS</td>
<td>5 mL</td>
<td>± 15%</td>
<td>500 mL</td>
</tr>
</tbody>
</table>

#### Table 11-2: Controls active in Respironics V60/V60 Plus ventilation modes

<table>
<thead>
<tr>
<th></th>
<th>CPAP</th>
<th>S/T</th>
<th>PCV</th>
<th>AVAPS</th>
<th>PPV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Timing</td>
<td></td>
<td>Rate*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>I-Time*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline pressure</td>
<td>CPAP</td>
<td>EPAP</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Inspiratory pressure</td>
<td></td>
<td>IPAP</td>
<td>Max P</td>
<td>Max P</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Min P</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rise Time</td>
<td></td>
<td>Rise*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>O₂</td>
<td>O₂</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Volume</td>
<td></td>
<td></td>
<td>VT</td>
<td>Max V</td>
<td></td>
</tr>
<tr>
<td>Ramp feature</td>
<td>Ramp Time</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mode-specific</td>
<td>C-Flex</td>
<td></td>
<td></td>
<td>PPV %</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Max E</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Max R</td>
<td></td>
</tr>
</tbody>
</table>

* Used in backup only
Technical specifications

**Patient data**

Table 11-3: Patient data ranges, resolutions, and accuracies during ventilation

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Range</th>
<th>Resolution</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Patient data window</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Breath phase/trigger indicator</td>
<td>Spont, Timed, Exhale</td>
<td>Color-coded display: Spont - turquoise, Timed - orange, Exhale - blue</td>
<td>N/A</td>
</tr>
<tr>
<td>PIP</td>
<td>0 to 50 cmH₂O</td>
<td>1 cmH₂O</td>
<td>± 2 cmH₂O</td>
</tr>
<tr>
<td>Pt. Leak</td>
<td>0 to 200 L/min BTPS</td>
<td>1 L/min</td>
<td>N/A</td>
</tr>
<tr>
<td>Pt. Trig</td>
<td>0 to 100%</td>
<td>1%</td>
<td>± 10%</td>
</tr>
<tr>
<td>Rate</td>
<td>0 to 90 BPM</td>
<td>1 BPM</td>
<td>± 1 BPM</td>
</tr>
<tr>
<td>T₁/T₅₀</td>
<td>0% to 91%</td>
<td>1%</td>
<td>± 5%</td>
</tr>
<tr>
<td>Tot.Leak</td>
<td>0 to 200 L/min BTPS</td>
<td>1 L/min</td>
<td>N/A</td>
</tr>
<tr>
<td>Vₑ</td>
<td>0 to 99.0 L/min BTPS</td>
<td>0.1 L/min</td>
<td>± 15% or 0.3 L/min (whichever is greater)</td>
</tr>
<tr>
<td>Vₜ</td>
<td>0 to 3500 mL BTPS</td>
<td>1 mL</td>
<td>± 15% for volumes above 200 mL Note: Accuracy specification was measured with patient circuit PN 582073</td>
</tr>
<tr>
<td><strong>Waveform window</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P waveform</td>
<td>0 to 50 cmH₂O</td>
<td>Time axis: 1 second</td>
<td>N/A</td>
</tr>
<tr>
<td>Vₑ waveform</td>
<td>-240 to 240 L/min BTPS</td>
<td>Time axis: 1 second</td>
<td>N/A</td>
</tr>
<tr>
<td>Vₜ waveform</td>
<td>0 to 3500 mL BTPS</td>
<td>Time axis: 1 second</td>
<td>N/A</td>
</tr>
</tbody>
</table>
Technical specifications

Alarms

Table 11-4 lists the adjustable alarm ranges and resolutions. Table 9-3 on page 9-7 describes other, nonadjustable alarms.

Table 11-4: Adjustable alarm ranges and resolutions

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Range</th>
<th>Resolution</th>
<th>Factory default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hi Rate (High Rate Alarm)</td>
<td>5 to 90 BPM</td>
<td>1 BPM</td>
<td>30 BPM</td>
</tr>
<tr>
<td>Lo Rate (Low Rate Alarm)</td>
<td>1 to 89 BPM</td>
<td>1 BPM</td>
<td>10 BPM</td>
</tr>
<tr>
<td>Hi VT (High Tidal Volume Alarm)</td>
<td>200 to 3500 mL BTPS</td>
<td>5 mL</td>
<td>2500 mL</td>
</tr>
<tr>
<td>Lo VT (Low Tidal Volume Alarm)</td>
<td>OFF, 5 to 1500 mL BTPS</td>
<td>5 mL</td>
<td>OFF</td>
</tr>
<tr>
<td>HIP (High Inspiratory Pressure Alarm)</td>
<td>5 to 50 cmH2O</td>
<td>1 cmH2O</td>
<td>50 cmH2O</td>
</tr>
<tr>
<td>LIP (Low Inspiratory Pressure Alarm)</td>
<td>OFF, 1 to 40 cmH2O</td>
<td>1 cmH2O</td>
<td>OFF</td>
</tr>
<tr>
<td>Lo Ve (Low Minute Ventilation Alarm)</td>
<td>OFF, 0.1 to 99.0 L/min BTPS</td>
<td>0.1 L/min</td>
<td>OFF</td>
</tr>
<tr>
<td>LIP T (Low Inspiratory Pressure Delay Time Alarm)</td>
<td>5 to 60 sec</td>
<td>1 sec</td>
<td>20 secs</td>
</tr>
</tbody>
</table>

Menu window settings

Table 11-5: Menu window settings and ranges

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brightness</td>
<td>1 to 5</td>
</tr>
<tr>
<td>Loudness</td>
<td>1 to 10</td>
</tr>
<tr>
<td>Mask/ET Selection</td>
<td>ET/Trach, 1, 2, 3, 4, Other</td>
</tr>
<tr>
<td>Exhalation Port Selection</td>
<td>• FEP (Philips Respironics Filtered Exhalation Port)</td>
</tr>
<tr>
<td></td>
<td>• DEP (Philips Respironics Disposable Exhalation Port)</td>
</tr>
<tr>
<td></td>
<td>• Whisper Swivel (Philips Respironics Whisper Swivel),</td>
</tr>
<tr>
<td></td>
<td>• PEV (Philips Respironics Plateau Exhalation Valve),</td>
</tr>
<tr>
<td></td>
<td>• Other (Other Exhalation Port),</td>
</tr>
<tr>
<td></td>
<td>• None (No circuit exhalation port)</td>
</tr>
<tr>
<td>Screen Lock</td>
<td>Off, On</td>
</tr>
<tr>
<td>Auto-Trak+ (optional)</td>
<td>Trigger: Normal, +1 to +7.</td>
</tr>
<tr>
<td></td>
<td>E-Cycle: -2 to -1, Normal, +1 to +6</td>
</tr>
</tbody>
</table>
## Diagnostic mode functions

Table 11-6: Diagnostic mode functions

<table>
<thead>
<tr>
<th>Function</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Language</td>
<td>English, Nederland, Français, Deutsch, Italiano, Português, Español, Dansk, Suomi, Norsk, Svenska, Chinês, Japanese, Türkçe</td>
</tr>
<tr>
<td>Date/Time</td>
<td>--</td>
</tr>
<tr>
<td>Pressure Units</td>
<td>cmH\textsubscript{2}O, hPa</td>
</tr>
<tr>
<td>Restore Default Settings</td>
<td>--</td>
</tr>
<tr>
<td>Software Options</td>
<td>--</td>
</tr>
<tr>
<td>Baud Rate</td>
<td>9,600, 19,200, 115,200</td>
</tr>
<tr>
<td>Alarm Volume Escalation(^*)</td>
<td>Enable, Disable (default)</td>
</tr>
<tr>
<td>Significant Event Log</td>
<td></td>
</tr>
<tr>
<td>Touch Screen Calibration</td>
<td></td>
</tr>
</tbody>
</table>

\(^*\) Available in Revision 2.30 software and above.
Technical specifications

Physical characteristics

Table 11-7: Physical characteristics

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dimensions</td>
<td>(33.7 cm) 13.3 in. (39.4 cm) 15.5 in. (42.9 cm) 16.5 in.</td>
</tr>
<tr>
<td>Mass (weight) of the V60/V60 Plus ventilator in its most usual configuration</td>
<td>12 kg (26 lb) with battery</td>
</tr>
<tr>
<td>Installed weight (V60/V60 Plus ventilator on stand, including accessories as listed)</td>
<td>V60/V60 Plus Ventilator with backup battery, ventilator stand, O₂ tank holder, two E-cylinders (full), two gauge clusters, oxygen analyzer with mount, humidifier with chamber, water bag and pole, nebulizer, circuit arm with mount, patient circuit with mask, O₂ manifold with hoses, remote alarm cable, and power cord. Weight: 64 kg (142 lb)</td>
</tr>
<tr>
<td>Maximum load (Mass/weight of the V60/V60 Plus ventilator stand, including its safe working load)</td>
<td>70.5 kg (155 lb)</td>
</tr>
</tbody>
</table>

Environmental specifications

Table 11-8: Environmental specifications

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature</td>
<td>Operating: 5 to 40 °C (41 to 104 °F) Storage/transport: -20 to 50 °C (-4 to 122 °F)</td>
</tr>
<tr>
<td>Relative humidity</td>
<td>Operating: 15 to 95% (noncondensing) Storage/transport: 10 to 95% relative (noncondensing)</td>
</tr>
<tr>
<td>Barometric pressure</td>
<td>Operating: 600 mmHg to 765 mmHg (80 kPa to 102 kPa): approximately -61 m to 1951 m (-200 ft to 6400 ft) relative to sea level Storage/transport: 450 mmHg to 765 mmHg (60 kPa to 102 kPa): approximately -61 m to 4267 m (-200 ft to 14000 ft) relative to sea level</td>
</tr>
</tbody>
</table>
## Pneumatic specifications

Table 11-9: Pneumatic specifications

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Specification</th>
</tr>
</thead>
</table>
| High-pressure oxygen supply                                              | Connector: DISS male, DISS female, NIST  
Pressure: 2.76 to 6.00 bar / 276 to 600 kPa / 40 to 87 psig  
Flow: 175 SLPM  
Connector: SIS  
Pressure: 3.31 to 6.00 bar / 331 to 600 kPa / 48 to 87 psig  
Flow: 175 SLPM |
| High-pressure oxygen supply (using V60/V60 Plus manifold)                | Connector: DISS male, DISS female, NIST  
Pressure: 3.10 to 6.00 bar / 310 to 600 kPa / 45 to 87 psig  
Flow: 175 SLPM  
Connector: SIS  
Pressure: 3.66 to 6.00 bar / 366 to 600 kPa / 53 to 87 psig  
Flow: 175 SLPM |
| Air supply                                                               | Integrated blower                                                                                                                                |
| Inspiratory outlet (to patient port)                                     | Connector: ISO 15 mm female/22 mm male conical                                                                                                 |
| Maximum limited pressure ($P_{LIMmax}$)                                  | 64 cmH₂O                                                                                                                                      |
| Maximum working pressure range ($P_{Wmax}$)                              | 5 to 50 cmH₂O, ensured by **High Inspiratory Pressure (HIP)** alarm limit  
None                                                                 |
| Subatmospheric pressures generated during exhalation                     | None                                                                                                                                              |
Technical specifications

Electrical specifications

Table 11-10: Electrical specifications

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td>AC voltage</td>
<td>100 to 240 VAC</td>
</tr>
<tr>
<td>AC frequency</td>
<td>50/60 Hz</td>
</tr>
<tr>
<td>AC power</td>
<td>300 VA</td>
</tr>
<tr>
<td>Battery</td>
<td>PN 1076374: 14.4 V, 11.0 Ah, 163 Wh</td>
</tr>
<tr>
<td></td>
<td>Maximum system current draw: 11 A</td>
</tr>
<tr>
<td></td>
<td>Charge voltage: +16.9 V maximum</td>
</tr>
<tr>
<td></td>
<td>Operating time: 360 minutes (6 hours) under normal conditions</td>
</tr>
</tbody>
</table>

NOTE: Any part that is added to the patient circuit changes the compliance and resistance of the ventilator breathing system. To achieve the ventilator performance specified in this manual, the ventilator breathing system must meet the specifications listed in Table 11-11.

Accessory requirements

Table 11-11: Accessory Requirements

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compliance</td>
<td>For volume accuracy requirements, the patient circuit compliance should be 0.98 mL/cmH₂O. For all other performance requirements except volume accuracy, the maximum compliance of the circuit can be up to 2.8 mL/cmH₂O.</td>
</tr>
<tr>
<td>Resistance</td>
<td>Maximum resistance of the breathing circuit and attachments: 2.9 cmH₂O at 60 L/min.</td>
</tr>
<tr>
<td>Bacteria filter</td>
<td>Dead Space: 66 ml Bacterial/Viral Filter Efficiency: &gt;99.99% Resistance (@ 0.5 L/s): 0.7 cmH₂O (hPa)/L/s Male Connector: 15 mm I.D./22 mm O.D. Female Connector: 22 mm I.D.</td>
</tr>
</tbody>
</table>
### Technical specifications

#### Alarm-related specifications

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Specification</th>
</tr>
</thead>
</table>
| Delay time from the onset of the alarm condition to the point that the representation of the alarm condition leaves the signal output port | <10 ms from onset of the alarm condition to the transmission of the signal  
<500 s until alarms received from the ventilator are displayed on the IntelliVue patient monitor. This delay is in addition to the alarm detection and processing delays of the external device.                                                                                                                                 |
| Remote alarm delay time                                                    | The time it takes the message to appear on the remote alarm depends on the characteristics of the device.                                                                                                                                                                                                                                     |
| Maximum time from the alarm condition triggering event to the generation of an alarm signal | All alarms except those listed below: < 10 s  
- Patient disconnect alarm: 11 s  
- Proximal pressure line disconnect alarm: 15 s  
- Low Inspiratory Pressure (LIP) alarm: user-configurable from 5 to 60 s  
- Check vent: Blower temperature high: 10 minutes                                                                                                                                                                                                                           |
| Mean delay, or the sum of mean delays, from the triggering event to the generation of an alarm signal | All alarms except those listed below: < 5 s  
- Cooling fan failure alarm: 10 s  
- Blower temperature alarm: 10 s  
- ADC wraparound: 6 s                                                                                                                                                                                                                                                         |
| Audio alarm loudness*                                                      | Highest volume setting: Average sound pressure level is approximately 76 dB(A)  
Lowest volume setting: Average sound pressure level is approximately 62 dB(A)                                                                                                                                                                                                                                                              |
| High-priority auditory alarm signal sound pressure level range, measured per ISO 3744:2010 ANSI/ | Average sound power level is approximately 54 dB(A) measured at the ventilator  
Average sound pressure level is approximately 46 dB(A) measured 1 m from the ventilator                                                                                                                                                                                                                                                |

* In accordance with 3rd Edition testing methods

#### Other specifications

<table>
<thead>
<tr>
<th>Parameter/Characteristic</th>
<th>Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Expected ventilator service life</td>
<td>10 years</td>
</tr>
<tr>
<td>Flow delivery</td>
<td>150 L/min at 40 cmH₂O at 1951 m (6400 ft) altitude (10% degradation in flow at 2286 m (7500 ft))</td>
</tr>
<tr>
<td>Flow range</td>
<td>0 to 240 L/min BTPS</td>
</tr>
</tbody>
</table>
### Technical specifications

#### Table 11-13: Other specifications (continued)

<table>
<thead>
<tr>
<th>Parameter/Characteristic</th>
<th>Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td>High flow therapy (3.00 software and above, and V60 Plus)</td>
<td>10 to 80 L/min BTPS</td>
</tr>
<tr>
<td><strong>NOTE:</strong> The maximum deliverable flow rate varies based on the orifice size of the nasal cannula and on the patient’s nasal passage resistance.</td>
<td></td>
</tr>
<tr>
<td>Pressure range</td>
<td>4 to 40 cmH₂O</td>
</tr>
<tr>
<td>Dynamic pressure regulation</td>
<td>± (2 cmH₂O + 4% of target)</td>
</tr>
<tr>
<td><strong>NOTE:</strong> Negative (subatmospheric) pressure settings are not available.</td>
<td></td>
</tr>
<tr>
<td>Start-up time</td>
<td>Ready to ventilate 9 seconds after power on</td>
</tr>
<tr>
<td>Triggering, cycling, and leak tolerance</td>
<td>As per the Digital Auto-Trak Sensitivity algorithms (see “Auto-Trak Sensitivity” on page 4-3)</td>
</tr>
</tbody>
</table>
| Inspiratory and expiratory pressure drop following equipment failure: measured at patient connection, when the recommended breathing system is in use. | < 2.0 cmH₂O (at 60 LPM)  
< 1.0 cmH₂O (at 30 LPM) |
| Maximum time required for the O₂ concentration in the delivered volume to change from a volume fraction of 21% to 90% using the worst-case ventilator breathing system or using the maximum internal volume ventilator breathing system. | The ventilator adjusts O₂ within one breath.  
FIO₂ within the gas delivery system and entire breathing circuit adjusts at the following rate:  
Up to 18 seconds for delivered volume of 500 mL  
Up to 17 seconds for delivered volume of 200 mL |
| The 10 s average input flow required by the ventilator for each gas at a pressure of 280 kPa | 107 SLPM |
| The maximum transient input flow averaged for 3 s required by the ventilator for each gas at a pressure of 280 kPa | 130 SLPM |
| Operational acoustics* | Average noise level is less than 45 dB(A) when measured 1 m from the ventilator |

* In accordance with 3rd Edition testing methods
Appendix A. First-time installation

WARNING: Philips-authorized personnel must install the ventilator. For information about installation, contact your Philips representative.

Before putting the ventilator into service for the first time, install it as described in this chapter.

Unpacking and inspection

Unpack the ventilator and inspect it for damage. Inspect the exterior cabinet of the ventilator for cracks, scratches, or blemishes. Inspect the front panel for scratches or abrasions. Correct and/or report any problems found to Philips before using the ventilator.

Before using the ventilator the first time, we recommend wiping the exterior clean and disinfecting components according to the instructions in Chapter 10.
First-time installation

Mounting the ventilator

CAUTION: To prevent possible damage to the ventilator, always secure it to its stand or securely place it on a flat, stable surface that is free of dirt and debris. Do not use the ventilator adjacent to, or stack it with, other equipment.

NOTE: If you mount the ventilator to a stand, make sure the stand is approved by Philips.

The ventilator may be mounted to the optional stand or placed on a flat, stable, clean surface. Figure A-2 shows the installed ventilator.

Align the ventilator above the stand as shown in Figure A-1. Make sure the four feet at the base of the ventilator are aligned to slide into the grooves at the top of the stand. Ensure that the locking lever on the back right of the stand locks into place.

Figure A-1: Mounting the V60/V60 Plus Ventilator on the stand
First-time installation

Figure A-2 shows the installed ventilator. Use the brakes to lock and unlock the wheels as needed. Make sure the wheels are unlocked before moving the ventilator.

Figure A-2: Respironics V60/V60 Plus Ventilator on stand
First-time installation

Installing the battery

WARNING: Installation or replacement of lithium batteries by inadequately trained personnel could result in a hazard.

WARNING: To reduce the risk of fire, explosion, leakage, or other hazard, take these precautions with respect to the battery:

- Do not attempt to disassemble, open, drop, crush, bend or deform, insert foreign objects into, puncture, or shred the battery pack; modify or remanufacture it; immerse or expose it to water or other liquids; expose it to fire, excessive heat (including soldering irons); or put it in a microwave oven.

- Replace the battery only with another battery specified by the manufacturer.

- Follow all instructions for proper use of the battery.

- Do not short-circuit the battery or allow metallic or conductive objects to contact the battery connector housing.

- Use the battery with the Respironics V60/V60 Plus Ventilator only.

Install the battery as follows (Figure A-3).

1. Shut down and then unplug the ventilator.

NOTE: Failure to properly shut down the ventilator before battery installation may result in erroneous alarms after power-on.

2. Remove the side panel by turning the captive fastener a ¼ turn and releasing.

3. Using a 3-mm hex wrench, remove the battery bracket by removing two screws.

4. Holding the battery so that the vent hole faces up and the Philips logo faces out, thread the battery cable through the battery bracket. Position and place the battery inside the battery compartment. Pinching the end of the battery connector, plug it in so that it locks in place.

5. Reinstall the battery bracket by replacing the two screws. Reinstall the side panel and secure the fastener with a ¼ turn clockwise.

6. Make sure the battery is properly installed by plugging the ventilator into an AC power receptacle and verifying that the yellow Battery (charged) LED on the front panel flashes. The flashing LED indicates the battery is being charged.
7. Attach the BATT label as shown in Figure 3-5 on page 3-9.

**WARNING:** Never attempt to disconnect or connect the battery during operation.

**CAUTION:** Following battery installation, if a Check Vent or Vent Inoperative alarm occurs when verifying ventilator operation, discontinue use of the ventilator immediately and contact Philips. The Vent Inoperative alarm occurs if AC power is disconnected and a battery is not installed, or if the battery is fully discharged.

**NOTE:** A new battery must be charged for at least 5 hours before being placed into service. Based on the age and state of the battery, it may take up to 16 hours or more to fully charge the battery.
First-time installation

Figure A-3: Installing the battery

Captive fastener
Bracket
Battery
Orient with vent hole on top and Philips logo facing out
Battery cable
First-time installation

Installing oxygen inlet connector and AC power cord

The Respironics V60/V60 Plus Ventilator destined for Japan, China, and the USA are typically pre-configured. V60/V60 Plus Ventilators shipped to other countries may require installation of the power cord and oxygen inlet connector.

1. Install the oxygen inlet connector as follows (Figure A-4):
   a. Gently fit connector into the hole provided with flat sides to the left and right.
   b. Install the oxygen inlet connector retaining plate. Tighten the two screws with a 2.5-mm hex wrench.

Figure A-4: Installing the oxygen inlet connector
First-time installation

**WARNING:** To prevent unintentional disconnection of the power cord, always use the correct, Philips-supplied power cord and lock it into place with the power cord retainer before you switch the ventilator on. The retainer is designed to hold the connector end of the Philips-supplied cord securely in place.

**WARNING:** The V60/V60 Plus Ventilator should not be positioned in a way that makes it difficult to disconnect from mains power if necessary. Disconnect from supply mains by removing the power cord from the wall outlet. The AC mains plug is used as disconnection device.

2. Secure the power cord with the power cord retainer (Figure A-5):
   a. Remove the power cord retainer by removing two screws.
   b. Connect the power cord that is appropriate to your region into the AC power connector.
   c. Reinstall the power cord retainer over the power cord, and tighten the screws with a 3.0-mm hex wrench.
First-time installation

Installing the oxygen manifold kit
If desired, install the oxygen manifold kit as described in the accompanying instructions.

Verifying ventilator operation and audible alarm
Perform the following steps to verify ventilator and audible alarm operation:

1. Assemble and install a patient circuit. (See Chapter 5 for instructions on installing a patient circuit.)
2. Power on the ventilator and verify that it completes the power-on self-test.
3. Disconnect the proximal pressure airway pressure line from the ventilator connector, and verify that the Proximal Pressure Line Disconnect alarm is annunciated (audio, visual, and flashing alarm LED).
4. Reconnect the proximal pressure line, and manually reset the alarm.
5. Turn the ventilator off.
6. Remove the patient circuit.

The ventilator is ready to be set up for use as described in Chapter 5.

Configuration and screen calibration
After completing the setup activities described in Chapter 5, set or check the ventilator settings for language, units of measure, and time in the diagnostic mode (see Appendix E). Calibrate the screen as required, referring to Appendix E.
First-time installation
Appendix B. Communications interface

WARNING: Connect to the ventilator only items that are specified as part of or compatible with the ventilator system. Additional equipment connected to medical electrical equipment must comply with the respective IEC or ISO standards. Furthermore, all configurations shall comply with the requirements for medical electrical systems (see IEC 60601-1-1 or clause 16 of edition 3 of IEC 60601-1, respectively). Anybody connecting additional equipment to medical electrical equipment configures a medical system and is therefore responsible for ensuring that the system complies with the requirements for medical electrical systems. Also be aware that local laws may take priority over the above mentioned requirements. If in doubt, consult Philips.

WARNING: The USB port is not currently available for use. DO NOT connect or attempt to power any equipment from the USB port.

WARNING: It is the responsibility of the end user to validate the compatibility and use of information transmitted from the ventilator to the device to be connected to the ventilator.

WARNING: The data provided through the communications interface is for reference only. Decisions for patient care should be based on the clinician’s observations of the patient.

The ventilator provides the following communications interface ports (Figure B-1):

- RS-232 serial port. Through this port the ventilator receives commands from a host computer or bedside monitoring system and responds with fixed-format records. The port is also used for ventilator servicing and software downloading.
- Remote alarm/nurse call port. Used to activate alarms remotely.
- USB and RJ-45 ethernet ports (not currently used)

Figure B-1: Location of communications interface ports
Communications interface

RS-232 serial port

The 25-pin D-sub RS-232 connector on the rear panel permits the ventilator to export parameters to a patient monitor or a hospital information system.

![Figure B-2: RS-232 serial connector pinout](image)

Supported communication protocols

The ventilator supports the following communication protocols:

- Philips IntelliBridge (VueLink): Use for connecting to Philips monitors (see "Using Philips IntelliBridge or VueLink" on page B-2).
- PVOI (Philips Ventilator Open Interface): Use for interfacing with non-Phillips monitors or devices. Contact your Philips representative to obtain the PVOI Developer’s Guide.
- Legacy (VRPT/SNDA): Fixed text-based protocol (not to be used for developing new driver interfaces). For this protocol specification, refer to the V60 Communications interface VRPT / SNDA Developers Guide (1147828MC), available at www.philips.com/IFU.

Using Philips IntelliBridge or VueLink

Using Philips monitors and the IntelliBridge or VueLink Open Interfaces

**NOTE:** Data displayed on the IntelliBridge or VueLink systems is for reference purposes only. Decisions for patient care should not be based solely on the data obtained through the IntelliBridge system.

The Respironics V60/V60 Plus Ventilator can communicate with a Philips patient monitor using the IntelliBridge Open Interface or the VueLink Open Interface.* Figure B-3 shows the required hardware setup. The IntelliBridge Open Interface and the VueLink Open Interface require a ventilator baud rate of 19,200. Check for the correct baud rate in the ventilator diagnostic mode (see “Baud Rate” on page E-10).

*NOTE: VueLink has been discontinued, but is still supported. Information is included for backwards compatibility only. The IntelliBridge EC10 is the VueLink Interface module successor.
Communications interface

Figure B-3: Connection from ventilator to Philips patient monitor

Data display

The data from your Respironics V60/V60 Plus Ventilator is displayed in several windows on your Philips monitor. This data may be labeled differently on the monitor than on the ventilator. Refer to Table B-1 to interpret these labels.

For more information, consult the documentation for your IntelliBridge or VueLink module and patient monitor.

Table B-1: Ventilator data displayed on Philips monitor

<table>
<thead>
<tr>
<th>Monitor label</th>
<th>Ventilator label</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Waveform</strong></td>
<td></td>
</tr>
<tr>
<td>AWP</td>
<td>P (airway pressure)</td>
</tr>
<tr>
<td>AWF</td>
<td>( V_E ) (flow)</td>
</tr>
<tr>
<td>AWV</td>
<td>V (volume)</td>
</tr>
<tr>
<td><strong>Monitored parameters</strong></td>
<td></td>
</tr>
<tr>
<td>%Bsp.t</td>
<td>Pt. Trigger</td>
</tr>
<tr>
<td>Leak</td>
<td>Pt. Leak or Tot.Leak</td>
</tr>
<tr>
<td>MINVOL</td>
<td>( V_E )</td>
</tr>
<tr>
<td>PIP</td>
<td>PIP</td>
</tr>
<tr>
<td>RRaw</td>
<td>Rate</td>
</tr>
<tr>
<td>Tin/Tt</td>
<td>( T_i/T_{TOT} )</td>
</tr>
<tr>
<td>TVexp</td>
<td>( V_T )</td>
</tr>
</tbody>
</table>
## Communications interface

### Table B-1: Ventilator data displayed on Philips monitor (continued)

<table>
<thead>
<tr>
<th>Monitor label</th>
<th>Ventilator label</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Modes</strong></td>
<td></td>
</tr>
<tr>
<td>Same as ventilator mode name</td>
<td>All modes except standby</td>
</tr>
<tr>
<td>STNDBY</td>
<td>Standby</td>
</tr>
<tr>
<td><strong>Control settings</strong></td>
<td></td>
</tr>
<tr>
<td>sEppv</td>
<td>Max E</td>
</tr>
<tr>
<td>Not shown</td>
<td>C-Flex</td>
</tr>
<tr>
<td>PAVsup</td>
<td>PPV %</td>
</tr>
<tr>
<td>sCPAP</td>
<td>CPAP</td>
</tr>
<tr>
<td>sEPAP</td>
<td>EPAP</td>
</tr>
<tr>
<td>sfgFl</td>
<td>Flow (in HFT)</td>
</tr>
<tr>
<td>sFI0₂</td>
<td>$O_2$</td>
</tr>
<tr>
<td>sInsTi</td>
<td>I-Time (Inspiratory Time)</td>
</tr>
<tr>
<td>sIPAP</td>
<td>IPAP</td>
</tr>
<tr>
<td>sPmax</td>
<td>Max P (AVAPS Maximum IPAP Pressure)</td>
</tr>
<tr>
<td></td>
<td>Max P (PPV Maximum IPAP Pressure)</td>
</tr>
<tr>
<td>sPmin</td>
<td>Min P (AVAPS Minimum IPAP Pressure)</td>
</tr>
<tr>
<td>sRisTi</td>
<td>Rise (Rise Time)</td>
</tr>
<tr>
<td>sRmpTi</td>
<td>Ramp Time</td>
</tr>
<tr>
<td>sRppv</td>
<td>Max R</td>
</tr>
<tr>
<td>sRRaw</td>
<td>Rate (Respiratory Rate)</td>
</tr>
<tr>
<td>sTV</td>
<td>$V_T$ (AVAPS Target Tidal Volume)</td>
</tr>
<tr>
<td>sVmax</td>
<td>Max V (PPV Maximum Volume Limit)</td>
</tr>
<tr>
<td>sVMode</td>
<td>Ventilation mode</td>
</tr>
</tbody>
</table>

### Alarm messages

<table>
<thead>
<tr>
<th>Alarm message</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>HIGH INSP PRESS</td>
<td>High Inspiratory Pressure</td>
</tr>
<tr>
<td>HIGH O2 SUPPLY</td>
<td>High $O_2$ Supply Pressure</td>
</tr>
<tr>
<td>HIGH RESP RATE</td>
<td>High Rate</td>
</tr>
<tr>
<td>HIGH EXH TV</td>
<td>High Tidal Volume</td>
</tr>
<tr>
<td>LOW INSP PRESS</td>
<td>Low Inspiratory Pressure</td>
</tr>
<tr>
<td>LOW BATTERY</td>
<td>Low Internal Battery</td>
</tr>
<tr>
<td>LOW FLOW</td>
<td>Cannot Reach Target Flow</td>
</tr>
<tr>
<td>LOW LEAK</td>
<td>Low Leak – CO₂ Rebreathing Risk</td>
</tr>
</tbody>
</table>
Table B-1: Ventilator data displayed on Philips monitor (continued)

<table>
<thead>
<tr>
<th>Monitor label</th>
<th>Ventilator label</th>
</tr>
</thead>
<tbody>
<tr>
<td>LOW EXH MV</td>
<td>Low Minute Ventilation</td>
</tr>
<tr>
<td>LOW O2 SUPPLY</td>
<td>Low O₂ Supply Pressure</td>
</tr>
<tr>
<td>LOW RESP RATE</td>
<td>Low Rate</td>
</tr>
<tr>
<td>LOW EXH TV</td>
<td>Low Tidal Volume</td>
</tr>
<tr>
<td>NO O2 SUPPLY</td>
<td>Oxygen Not Available</td>
</tr>
<tr>
<td>OCCLUSION</td>
<td>Patient Circuit Occluded</td>
</tr>
<tr>
<td>PT. DISCONNECT</td>
<td>Patient Disconnect</td>
</tr>
<tr>
<td>PPV MAX P</td>
<td>PPV Max P</td>
</tr>
<tr>
<td>PPV MAX V</td>
<td>PPV Max V</td>
</tr>
<tr>
<td>PRESS REG HIGH</td>
<td>Pressure Regulation High</td>
</tr>
<tr>
<td>PROX DISCONNECT</td>
<td>Proximal Pressure Line Disconnect</td>
</tr>
<tr>
<td>Vent CHK DEVICE</td>
<td>Check Vent:</td>
</tr>
<tr>
<td>VENT ON BATTERY</td>
<td>Running on Internal Battery</td>
</tr>
<tr>
<td>Ventilation parameters blanked</td>
<td>Vent Inoperative xxxx</td>
</tr>
</tbody>
</table>

WARNING: To prevent possible patient injury due to nonannunciating alarms, verify the operation of any remote alarm device before use.

WARNING: To ensure the functionality of the remote alarm, connect only Respironics-approved cables to the remote alarm port.

CAUTION: The remote alarm port is intended to connect only to an SELV (safety extra-low voltage and ungrounded system with basic insulation to ground), in accordance with IEC 60601-1. To prevent damage to the remote alarm, make sure the signal input does not exceed the maximum rating of 24 VAC or 36 VDC at 500 mA with a minimum current of 1 mA.

NOTE: Selecting Alarm Silence deactivates the remote alarm.

The remote alarm (nurse call) port allows ventilator alarm conditions to be annunciated at locations away from the ventilator (for example, when the ventilator is in an isolation room). The ventilator sends alarm signals to a remote alarm through the connector at the rear of the ventilator (Figure B-1 on page B-1). Figure B-4 shows the pin assignments for this connector. The connector is a standard ¼-inch, female, audio (ring, tip, sleeve) connector.

The ventilator signals an alarm using either a normally open (NO) or normally closed (NC) relay contact. The de-energized state of the relay represents an
Communications interface

alarm state (any high-priority alarm) and the energized state represents a non-alarm state. This application requires one of the cables listed in Table B-2.

Figure B-4: Remote alarm port

Table B-2: Remote alarm cable kits

<table>
<thead>
<tr>
<th>Description</th>
<th>System</th>
<th>Part Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Remote alarm cable kit, alarm state = open</td>
<td>Use on Normally Closed nurse call systems (system expects to see open contacts when ventilator alarms). For ¼” jack.</td>
<td>1003741</td>
</tr>
<tr>
<td>Remote alarm cable kit, alarm state = closed</td>
<td>Use on Normally Opened nurse call systems (system expects to see closed contacts when ventilator alarms). For ¼” jack.</td>
<td>1003742</td>
</tr>
<tr>
<td>Remote alarm cable kit</td>
<td>Philips Respironics (LifeCare)</td>
<td>1003743</td>
</tr>
</tbody>
</table>
Appendix C. Parts and accessories

This appendix lists parts and accessories supplied by Philips that are compatible with the Respironics V60/V60 Plus Ventilator. All parts and accessories are not available in all markets.

**WARNING:** Avoid adding resistive circuit components on the patient side of the proximal pressure line. Such components may defeat the disconnect alarm.

**NOTE:** Resistive components can include but are not limited to HMEs, proximal flow sensors, a filter at the patient connection, or a narrow diameter circuit attached to a mask.

**NOTE:** To ensure the correct performance of the ventilator and the accuracy of patient data, we recommend you use only Respironics-approved accessories with the ventilator.

For the most current list of approved accessories plus comprehensive ordering information for compatible parts available from Philips, contact your Philips representative or refer to the V60/V60 Plus ordering and accessories guide (downloadable from philips.com/ifu or philips.com/hrcmanuals).
NOTE: Pediatric masks intended for patients weighing less than 20 kg (44 lb) are not approved for use with the V60/V60 Plus Ventilator.

**Ventilation interfaces**

Compatible interfaces include entrainment elbow (EE) versions from these product lines:

<table>
<thead>
<tr>
<th>Description</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Respironics Contour Deluxe</td>
<td>Nasal</td>
</tr>
<tr>
<td>Respironics AP111</td>
<td></td>
</tr>
<tr>
<td>Respironics PerformaTrak</td>
<td>Oro-nasal</td>
</tr>
<tr>
<td>Respironics AF541, over-the-nose</td>
<td></td>
</tr>
<tr>
<td>Respironics AF541, under-the-nose</td>
<td></td>
</tr>
<tr>
<td>Respironics AF531</td>
<td></td>
</tr>
<tr>
<td>Respironics PerforMax, adult only</td>
<td>Total</td>
</tr>
</tbody>
</table>

**HFT interfaces**

For use with V60 Plus ventilators and V60 ventilators that have 3.00 software (or higher) and the HFT option installed.

<table>
<thead>
<tr>
<th>Description</th>
<th>Part number</th>
<th>Order number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Respironics AC611 high flow nasal cannula</td>
<td>1128903</td>
<td>989805654621</td>
</tr>
<tr>
<td>Respironics AC611 high flow nasal cannula with filtered exhalation port (FEP) connector</td>
<td>1130625</td>
<td>989805655111</td>
</tr>
</tbody>
</table>

**O₂ analyzer/monitor**

<table>
<thead>
<tr>
<th>Description</th>
<th>Part number</th>
<th>Order number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Analytical Industries All-2000M oxygen analyzer/monitor</td>
<td>1129064</td>
<td>989805654731</td>
</tr>
<tr>
<td>O₂ sensor, All-2000M</td>
<td>1130625</td>
<td>989805655111</td>
</tr>
<tr>
<td>Tee adapter, O₂ sensor</td>
<td>1128903</td>
<td>989805654621</td>
</tr>
<tr>
<td>Tee adapter, O₂ sensor, threaded, amber</td>
<td>1020380</td>
<td>453561509031</td>
</tr>
</tbody>
</table>
## Patient breathing circuits

<table>
<thead>
<tr>
<th>Description</th>
<th>Quantity</th>
<th>Part number</th>
<th>Order number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Circuits Supporting Noninvasive Ventilation and High-Flow Therapy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patient circuit, single-limb, heated, proximal pressure line, single-use, adult (Fisher &amp; Paykel RT139).</td>
<td>10</td>
<td>1020523</td>
<td>989805610851</td>
</tr>
<tr>
<td>Patient circuit, single-limb, heated, filtered exhalation port (FEP), humidifier chamber, proximal pressure line, single-use, 22-mm ID, WILAmed (Not available in North America)</td>
<td>10</td>
<td>1122059</td>
<td>989805653191</td>
</tr>
<tr>
<td>Patient circuit, single-limb, heated, filtered exhalation port (FEP), humidifier chamber, proximal pressure line, single-use, 22-mm ID, Fisher &amp; Paykel RT239</td>
<td>10</td>
<td>1135739</td>
<td>989805658961</td>
</tr>
<tr>
<td>Circuits Supporting Noninvasive Ventilation Only</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patient circuit, single-limb, non-heated, filtered exhalation port (FEP), proximal pressure line, single-use, 22-mm ID</td>
<td>10</td>
<td>1065830 (with inspiratory and exhalation port filters)</td>
<td>989805621311</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1065832 (with inspiratory port filter only)</td>
<td>989805621321</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1069210 (with no filters)</td>
<td>989805634871</td>
</tr>
<tr>
<td>Patient circuit, single-limb, non-heated, disposable exhalation port (DEP), proximal pressure line, single-use, 22-mm ID</td>
<td>10</td>
<td>582073</td>
<td>989805609611</td>
</tr>
<tr>
<td>Patient circuit, single-limb, non-heated, filtered exhalation port (FEP), water trap, proximal pressure line, single-use, 22-mm ID</td>
<td>10</td>
<td>652002</td>
<td>989805609681</td>
</tr>
</tbody>
</table>

## Humidifiers

<table>
<thead>
<tr>
<th>Description</th>
<th>Quantity</th>
<th>Part number</th>
<th>Order number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Humidifiers</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Respiratory humidifier, Fisher &amp; Paykel MR850</td>
<td>---</td>
<td>---</td>
<td>Refer to the V60 ordering and accessories guide</td>
</tr>
<tr>
<td>Respiratory humidifier, WILAmed AIRcon Gen2, 230V (Not available in North America)</td>
<td>---</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td>Chambers</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Humidifier chamber, adult, Fisher &amp; Paykel MR290</td>
<td>10</td>
<td>22104</td>
<td>989805642931</td>
</tr>
<tr>
<td>Humidifier chamber, autofill, WILAmed (Not available in North America)</td>
<td>30</td>
<td>1121825</td>
<td>989805653111</td>
</tr>
</tbody>
</table>
### Parts and accessories

#### Bacteria filters

<table>
<thead>
<tr>
<th>Description</th>
<th>Quantity</th>
<th>Part number</th>
<th>Order number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Filter inspiratory, 22-mm male x 22-mm female, single-use</td>
<td>10</td>
<td>342077</td>
<td>989805609521</td>
</tr>
<tr>
<td>Filter, patient pressure, single-use</td>
<td>1</td>
<td>1002362</td>
<td>453561517101</td>
</tr>
</tbody>
</table>

#### Nebulizers

<table>
<thead>
<tr>
<th>Description</th>
<th>Quantity</th>
<th>Part number</th>
<th>Order number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aerogen Solo nebulizer (for inline applications)</td>
<td>---</td>
<td>---</td>
<td>Refer to the V60 ordering and accessories guide</td>
</tr>
<tr>
<td>NIVO generator (for mask elbow applications)</td>
<td>5</td>
<td>1076302</td>
<td>989805634301</td>
</tr>
</tbody>
</table>

#### Operator maintenance parts

<table>
<thead>
<tr>
<th>Description</th>
<th>Quantity</th>
<th>Part number</th>
<th>Order number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cooling fan filter</td>
<td>5</td>
<td>1054280</td>
<td>453561507301</td>
</tr>
<tr>
<td>Air inlet filter</td>
<td>5</td>
<td>1054279</td>
<td>453561505991</td>
</tr>
</tbody>
</table>

#### Other parts

Compatible with all V60/V60 Plus Ventilators:

<table>
<thead>
<tr>
<th>Description</th>
<th>Part number</th>
<th>Order number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Test lung, hard-sided (low-compliance)</td>
<td>1021671</td>
<td>989805611871</td>
</tr>
<tr>
<td>Backup battery</td>
<td>1076374</td>
<td>989805626941</td>
</tr>
</tbody>
</table>
Appendix D. Regulatory compliance

**Electromagnetic compatibility (EMC)**

| IEC 60601-1-2; 2014, Ed. 4.0 | Medical electrical equipment - Part 1-2: General requirements for basic safety and essential performance - Collateral Standard: Electromagnetic disturbances |

Medical equipment needs special precautions regarding electromagnetic compatibility (EMC) and needs to be installed and put into service according to the EMC information provided in this document.

**WARNING:** The V60/V60 Plus Ventilator may cause radio interference or may disrupt the operation of nearby equipment. It may be necessary to take mitigation measures, such as re-orienting or relocating the ventilator or shielding the location.

**WARNING:** Use of non-approved accessories, transducers or cables may increase EMC emissions or decrease the EMC immunity performance of the equipment.

**WARNING:** Use portable radio-frequency (RF) communications equipment (including peripherals such as antenna cables and external antennas) no closer than 30 cm (12 in.) to any part of the ventilator system, including cables specified for use with the ventilator. Otherwise, equipment performance can be degraded. If higher immunity test levels than those specified in IEC 60601-1-2:2014, Table 9, are used, the minimum separation distance may be lowered. Lower minimum separation distances shall be calculated using the equation specified in chapter 8.10 of the standard.

**WARNING:** Do not use the ventilator near radio-frequency identification (RFID) or electromagnetic security systems. The ventilator may disrupt the operation of this equipment.

**WARNING:** This equipment is designed to comply with IEC 60601-1-2. This equipment generates, uses, and can radiate radio-frequency energy and, if not installed and used in accordance with instructions, may cause harmful interference to other devices in the vicinity. However, there is no guarantee that interference will not occur in a particular installation. Harmful interference to other devices can be determined by turning this equipment on and off. Try to correct the interference using one or more of the following:

- Reorient or relocate the receiving device.
- Increase the separation between the equipment.
Regulatory compliance

- Connect the equipment to an outlet on a different circuit from that to which the other devices are connected and contact Philips Service for help.

**WARNING:** Take care when operating the ventilator around other equipment, to avoid reciprocal interference. Potential electromagnetic interference (EMI), electrostatic discharge (ESD), or other interference can occur to the ventilator or other equipment. Interference can interrupt ventilator operation or degrade system performance, which can result in patient injury. Try to minimize this interference by not using other equipment in conjunction with the ventilator. If adjacent or stacked use is necessary, follow the recommendations for placement of the equipment in “Electromagnetic compatibility declaration” on page D-1. Monitor the ventilator and nearby devices to verify normal operation.

**WARNING:** Do not use the ventilator near active high frequency (HF) surgical equipment, medical devices such as X-ray devices and diathermy, or in an RF-shielded room of medical equipment or system for magnetic resonance imaging, where the intensity of electromagnetic (EM) disturbances is high. These devices may degrade the performance of the ventilator.

Electromagnetic emissions

NOTE: The emissions characteristics of this equipment make it suitable for use in industrial areas and hospitals (CISPR 11 class A).

<table>
<thead>
<tr>
<th>Guidance and manufacturer's declaration - electromagnetic emissions</th>
</tr>
</thead>
<tbody>
<tr>
<td>The V60/V60 Plus Ventilator is intended for use in the electromagnetic environment specified below. The user of the V60/V60 Plus Ventilator should assure that it is used in such an environment.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Emissions test</th>
<th>Compliance</th>
<th>Electromagnetic enforcement - guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF Emissions CIS-PR 11</td>
<td>Group 1</td>
<td>The V60/V60 Plus Ventilator uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause interference in nearby electronic equipment.</td>
</tr>
<tr>
<td>RF Emissions CIS-PR 11</td>
<td>Class A</td>
<td>The V60/V60 Plus Ventilator is suitable for use in all establishments, except for those that are domestic or directly connected to the public low voltage power supply network that supplies buildings used for domestic purposes.</td>
</tr>
<tr>
<td>Harmonic emissions IEC 61000-3-2</td>
<td>Class A</td>
<td></td>
</tr>
<tr>
<td>Voltage fluctuations/flicker emissions IEC 61000-3-3</td>
<td>Complies</td>
<td></td>
</tr>
</tbody>
</table>
# Regulatory compliance

## Electromagnetic immunity

### Guidance and manufacturer’s declaration - electromagnetic immunity

The V60/V60 Plus Ventilator is intended for use in the electromagnetic environment specified below. The user of the V60/V60 Plus Ventilator should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity test</th>
<th>IEC 60601-1-2 test level</th>
<th>Compliance level</th>
<th>Electromagnetic environment guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrostatic discharge (ESD)</td>
<td>±8 kV contact</td>
<td>±8 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>±15 kV air</td>
<td>±15 kV air</td>
<td></td>
</tr>
<tr>
<td>Electrical fast transient/burst</td>
<td>±2 kV for power supply lines.</td>
<td>±2 kV for power supply lines.</td>
<td>Mains power quality should be that of a typical hospital environment.</td>
</tr>
<tr>
<td>IEC 61000-4-4</td>
<td>±1 kV for input / output lines</td>
<td>±1 kV for input / output lines</td>
<td></td>
</tr>
<tr>
<td>Surge</td>
<td>±1 kV differential mode</td>
<td>±1 kV differential mode</td>
<td>Mains power quality should be that of a typical hospital environment.</td>
</tr>
<tr>
<td>IEC 61000-4-5</td>
<td>±2 kV common mode</td>
<td>±2 kV common mode</td>
<td></td>
</tr>
<tr>
<td>Voltage dips, short interruptions and voltage variations on power supply input lines</td>
<td>0% $U_T$ for 0.5 cycles 0% $U_T$ for 1.0 cycle 70% $U_T$ for 25 cycles (50 Hz)/30 cycles (60 Hz) 0% $U_T$ for 250 cycles (50 Hz)/300 cycles (60 Hz)</td>
<td>0% $U_T$ for 0.5 cycles 0% $U_T$ for 1.0 cycle 70% $U_T$ for 25 cycles (50 Hz)/30 cycles (60 Hz) 0% $U_T$ for 250 cycles (50 Hz)/300 cycles (60 Hz)</td>
<td>Mains power quality should be that of a typical hospital environment. If the user of the V60/V60 Plus Ventilator requires continued operation during power mains interruptions, it is recommended that the V60/V60 Plus Ventilator be powered from an uninterruptible power supply or a battery.</td>
</tr>
<tr>
<td>IEC 61000-4-11</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Power frequency (50/60 Hz) magnetic field</td>
<td>30 A/m</td>
<td>30 A/m</td>
<td>Power frequency magnetic fields should be at levels characteristic of a typical location in a typical 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.
Regulatory compliance

**Guidance and manufacturer's declaration - electromagnetic immunity**

The V60/V60 Plus Ventilator is intended for use in the electromagnetic environment specified below. The customer or the user of the V60/V60 Plus Ventilator should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity test</th>
<th>IEC 60601-1-2 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><strong>Portable and mobile RF communications equipment should be used no closer to any part of the V60/V60 Plus Ventilator, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.</strong> Recommended separation distance:</td>
</tr>
<tr>
<td></td>
<td>3 Vrms 150 kHz to 80 MHz outside ISM bands(^a)</td>
<td>3 Vrms</td>
<td></td>
</tr>
<tr>
<td></td>
<td>6 Vrms 150 kHz to 80 MHz in ISM bands(^a)</td>
<td>6 Vrms</td>
<td></td>
</tr>
<tr>
<td>Radiated RF</td>
<td>IEC 61000-4-3</td>
<td>3 V/m 80 MHz to 2.7 GHz</td>
<td>80 MHz to 800 MHz, where ( E_1 = 3 \text{ V/m} )</td>
</tr>
<tr>
<td></td>
<td>380 MHz to 390 MHz</td>
<td>27 V/m</td>
<td></td>
</tr>
<tr>
<td></td>
<td>430 MHz to 470 MHz</td>
<td>28 V/m</td>
<td></td>
</tr>
<tr>
<td></td>
<td>704 MHz to 787 MHz</td>
<td>9 V/m</td>
<td></td>
</tr>
<tr>
<td></td>
<td>800 MHz to 960 MHz</td>
<td>28 V/m</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1.700 GHz to 1.990 GHz</td>
<td>28 V/m</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2.400 GHz to 2.570 GHz</td>
<td>9 V/m</td>
<td></td>
</tr>
<tr>
<td></td>
<td>5.100 GHz to 5.800 GHz</td>
<td>9 V/m</td>
<td></td>
</tr>
<tr>
<td></td>
<td>800 MHz to 2.7 GHz</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>80 MHz to 2.7 GHz</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2.400 GHz to 2.570 GHz</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>5.100 GHz to 5.800 GHz</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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

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

\( d = \frac{\frac{3.5}{V_1}}{P} \)  
where \( V_1 = 3 \text{ Vrms} \)

\( d = \frac{\frac{12}{V_2}}{P} \)  
where \( V_2 = 6 \text{ Vrms} \)

\( d = \frac{\frac{12}{E_1}}{P} \)  
80 MHz to 800 MHz, where \( E_1 = 3 \text{ V/m} \)

\( d = \frac{\frac{23}{E_1}}{P} \)  
800 MHz to 2.7 GHz, where \( E_1 = 3 \text{ V/m} \)

---

\(^a\) 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 2.7 GHz are intended to decrease the likelihood that mobile/ portable communications equipment could cause interference if it is inadvertently brought into patient areas. For this reason, an additional factor of 10/3 has been incorporated into the formulae used in calculating the recommended separation distance in meters (m).

\(^c\) Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, should be less than the compliance level in each frequency range.

\(^d\) Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m.

\(^e\) According to IEC 60601-1-2: 2014
### Recommended separation distances between portable and mobile RF communications equipment and the V60/V60 Plus Ventilator

<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>150 kHz to 80 MHz outside ISM bands</td>
<td>150 kHz to 80 MHz in ISM bands</td>
</tr>
<tr>
<td>$d = \left[ \frac{3.5}{P} \right]^{1/2}$</td>
<td>$d = \left[ \frac{12}{E_1} \right]^{1/2}P$</td>
</tr>
<tr>
<td>0.01</td>
<td>0.12</td>
</tr>
<tr>
<td>0.1</td>
<td>0.37</td>
</tr>
<tr>
<td>1</td>
<td>1.17</td>
</tr>
<tr>
<td>10</td>
<td>3.69</td>
</tr>
<tr>
<td>100</td>
<td>11.67</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 determined 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:** 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. **NOTE 3:** An additional factor of 10/3 has been incorporated into the formulae 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 2.7 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. **NOTE 5:** The minimum separation distance for RF communication equipment operating within the following frequency bands is 0.3 m:

- 380 - 390 MHz (TETRA 400)
- 430 - 470 MHz (GMRS 460, FRS 460)
- 704 - 787 MHz (LTE Band 13, 17)
- 800 - 960 MHz (GSM 800/900, TETRA 800, iDEN 820, CDMA 850, LTE Band 5)
- 1 700 - 1 990 MHz (GSM 1800; CDMA 1900; GSM 1900; DECT; LTE Band 1, 3, 4, 25; UMTS)
- 2 400 - 2 570 MHz (Bluetooth, WLAN, 802.11 b/g/n, RFID 2450, LTE Band 7)
- 5 100 - 5 800 MHz (WLAN 802.11 a/n)
**Regulatory compliance**

**RF immunity**

<table>
<thead>
<tr>
<th>RFID specification</th>
<th>Test Frequency</th>
<th>Test level (RMS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ISO 14223</td>
<td>134.2 kHz</td>
<td>65 A/m</td>
</tr>
<tr>
<td>ISO/IEC 14443-3 (Type A)</td>
<td>13.56 MHz</td>
<td>7.5 A/m</td>
</tr>
<tr>
<td>ISO/IEC 14443-4 (Type B)</td>
<td>13.56 MHz</td>
<td>7.5 A/m</td>
</tr>
<tr>
<td>ISO/IEC 15693 (ISO 18000-3 Mode 1)</td>
<td>13.56 MHz</td>
<td>5 A/m</td>
</tr>
<tr>
<td>ISO 18000-3 Mode 3</td>
<td>13.56 MHz</td>
<td>12 A/m</td>
</tr>
<tr>
<td>ISO/IEC 18000-7</td>
<td>433 MHz</td>
<td>3 V/m</td>
</tr>
<tr>
<td>ISO/IEC 18000-63 Type C</td>
<td>860-960 MHz</td>
<td>54 V/m</td>
</tr>
<tr>
<td>ISO/IEC 18000-4 Mode 1</td>
<td>2.45 GHz</td>
<td>54 V/m</td>
</tr>
</tbody>
</table>

**RFID reader separation distance**

The testing specified in this standard applies to medical electrical equipment and systems used near RFID readers. A medical electrical equipment or system that is in conformity with this standard is qualified for use as close as 2.5 cm for some RFID applications and 20 cm for others, based on the maximum amount of radio frequency (RF) output power allowed by the Federal Communications Commission (FCC) and typical use distances. Medical electrical equipment and systems that meet less stringent standards, such as IEC 60601-1-2, could be used safely with RFID readers if the RF output power of the RFID reader is less than the allowed maximum or it can be assured that the separation distance between the RFID Reader and the medical electrical equipment or system will always be much greater than 20 cm.

Guidance on EMC usually includes the parameters of the RF output of the source (P), the RF immunity of the medical electrical equipment or system, and the distance between them (d). In general, a medical electrical equipment or system should operate safely when it experiences field strength (E) that is equal to or lower than its specified RF immunity. In general, the field strength falls off proportional to 1/d, although care must be taken because some environments (where large metallic objects such as HVAC ducts and cabinetry are present) can cause reflections such that field strength in certain close-in ranges can be higher further away from the medical electrical equipment or system that is at closer distances.
So, in addition to selection of RFID systems and medical electrical equipment and systems, EMC management includes maintaining adequate separation distances between them.

For medical electrical equipment and systems that meet this standard, the minimum necessary separation distances are shown in Annex J of the AIMS standard 7351731, as well as the guidance required by standards such as IEC 60601-1-2.

For medical electrical equipment and systems that do not meet this standard, the minimum necessary separation distances can be determined using the guidance required by standards such as IEC 60601-1-2.

In general, the recommended minimum separation distance \( d \) is of the form:

\[
d = \left[ \frac{k}{E} \right] \sqrt{P}
\]

where \( E \) is the RF immunity of the medical electrical equipment or system, \( P \) is the maximum rated RF output power of the RFID system, and \( k \) is an antenna efficiency factor. In Edition 3 of IEC 60601-1-2 \[23\] and earlier editions, \( k \) was assumed to be 3.5 for frequencies less than 800 MHz and 7 at frequencies greater than or equal to 800 MHz. In Edition 4 \[24\], \( k \) is assumed to be 6 for all frequencies.

### Cables That May Affect IEC 60601-1-2 Compliance

Cables, such as remote alarm cables, longer than 3.05-m may affect the compliance of the ventilator system with the emissions and immunity requirements of IEC 60601-1-2.

### WEEE recycling directive

Waste electrical and electronic equipment (WEEE) recycling directive.

Waste electrical and electronic equipment must not be disposed of as unsorted municipal waste at the end of its expected service life. It must be collected separately and must be disposed of per local regulations. Contact your Philips authorized representative for information concerning the decommissioning of your equipment.

Compliant with the WEEE recycling directive.

If you are subject to the WEEE directive, refer to www.philips.com/recycling for the passport for recycling this product.

Dispose of this device in accordance with local collections and recycling regulations. For more information, visit www.philips.com/recycling.

This device contains a lithium ion battery that cannot be recycled. Please contact customer service for safe disposal instructions of this part.
## Regulatory compliance

### Classification

<table>
<thead>
<tr>
<th>Protection Against Electric Shock</th>
<th>Class I</th>
</tr>
</thead>
<tbody>
<tr>
<td>Degree of Protection Against Electric Shock</td>
<td>Type B</td>
</tr>
<tr>
<td>Degree of protection against ingress of particulate matter and water given by the enclosure</td>
<td>IP21 Protected against solid objects ≥ 12.5 mm in diameter. Protected against vertically falling water drops when enclosure is tilted up to 15°</td>
</tr>
<tr>
<td>Mode of operation</td>
<td>Continuous Operation</td>
</tr>
<tr>
<td>Method of sterilization</td>
<td>Not intended to be sterilized</td>
</tr>
<tr>
<td>Suitability for use in an O₂-rich environment</td>
<td>Not suitable</td>
</tr>
</tbody>
</table>

### Safety

<table>
<thead>
<tr>
<th>Standard</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>IEC 60601-1; 2012, Ed. 3.1</td>
<td>Medical electrical equipment – Part 1: General requirements for basic safety and essential performance</td>
</tr>
<tr>
<td>IEC 60601-1-6; 2013, Ed. 3.1</td>
<td>Medical electrical equipment – Part 1-6: General requirements for basic safety and essential performance; collateral standard: usability</td>
</tr>
<tr>
<td>IEC 60601-1-8; 2012, Ed. 2.1</td>
<td>Medical electrical equipment – Part 1-8: General requirements for basic safety and essential requirements; collateral standard - alarm systems</td>
</tr>
<tr>
<td>IEC 62366-1, 2015, Ed. 1.1</td>
<td>Medical devices - Application of usability engineering to medical devices</td>
</tr>
<tr>
<td>ISO 14971; 2007</td>
<td>Medical devices - Application of risk management to medical devices</td>
</tr>
<tr>
<td>EN ISO 14971; 2012</td>
<td>Medical devices - Application of risk management to medical devices</td>
</tr>
<tr>
<td>ISO 80601-2-12; 2011</td>
<td>Medical electrical equipment – Particular requirements for basic safety and essential performance of critical care ventilators</td>
</tr>
<tr>
<td>IEC 60529; 2013, Ed. 2.2</td>
<td>Degrees of protection provided by enclosures (IP Code)</td>
</tr>
<tr>
<td>IEC 62304; 2015, Ed. 1.1</td>
<td>Medical device software - Software life cycle processes</td>
</tr>
</tbody>
</table>
Regulatory compliance

**Applied parts**

The V60/V60 Plus Ventilator system includes these applied parts:
- Patient interfaces
- Tracheal and endotracheal tubes
- High flow therapy nasal cannula and interface
- Nebulizer T-adapter
- Ventilator breathing system - Type B (inspiratory or expiratory pathways through which gas flows at respiratory pressures and bounded by the ports through which gas enters and the patient-connection port), which can also include the following:
  - Humidifier chamber
  - Patient circuits

**Accessible parts**

The V60/V60 Plus Ventilator system includes these accessible parts:
- Filters
- Roll stand
- Humidifier
- O₂ monitor/analyzer
- Nebulizer controller

**Detachable components**

The V60/V60 Plus Ventilator system includes these detachable components (see Figure 3-4 and Figure 3-5):
- Left side panel
- Right side panel
- Air inlet filter (under side panel)

**Essential performance**

Per ISO/EN 80601-2-12: 2011, Medical electrical equipment Part 2-12: Particular requirements for basic safety and essential performance of critical care ventilators, the ventilator’s essential performance requirements are given in “Control settings” on page 11-1, “Alarms and other messages” on page 9-7, and “Table of monitored parameters” on page 8-2. Alarms, including O₂ supply failure alarms and gas failure alarms, are identified in “Alarms and other messages” on page 9-7. AC mains power information is given in “Connecting to AC power” on page 5-2. Battery backup information is given in “About the backup battery” on page 3-10. Gas connection information is given in “Connecting oxygen” on page 5-1 and “Installing oxygen inlet connector and AC power cord” on page A-7.
Regulatory compliance
Appendix E. Diagnostic mode

In the diagnostic mode you select the language of software display, set the date and time, select pressure units, enable software options, and calibrate the touchscreen.

NOTE: The diagnostic mode is primarily for use by authorized service personnel to download software and perform other diagnostic procedures.

Entering the diagnostic mode

WARNING: To prevent possible patient injury, do not enter the diagnostic mode while a patient is connected to the ventilator. Verify that the patient is disconnected before proceeding.

Enter the diagnostic mode as follows:

1. Make sure the patient is disconnected and the ventilator is powered off.

2. Press and hold the Accept button on the top-right front of the ventilator and turn on the ventilator by pressing the ON/Shutdown key.

The screen displays “Press \( \square \) again for Diagnostics or wait for Ventilation.”
Diagnostic mode

3. Within less than 5 seconds, release and press the Accept button again. The Diagnostics Menu (Figure E-1) is displayed.

   ![Diagnostics Menu](image)

   *If buttons are unresponsive, press *key* to calibrate touch screen.

   **WARNING**
   The Diagnostics Mode is not to be used when a patient is connected to the ventilator. Verify that the patient is disconnected prior to proceeding.

   **Figure E-1: Diagnostics Menu**

4. Select the desired function.
System settings

From the System Settings screen (Figure E-2) you can perform the functions below.

![System Settings screen](image)

Figure E-2: System Settings screen
Diagnosis mode

Language
The Language function lets you set the language of software display.

1. From the System Settings screen, select Language to display the Set Language screen (Figure E-3).

![Set Language screen](image)

Figure E-3: Set Language screen 1
2. The active language is shown in white type. Select the new language.

3. A second Set Language screen is displayed (Figure E-4). Select Ventilator Shutdown to apply the change. The change is effective after you restart the ventilator.

![Set Language Screen](image)

Figure E-4: Set Language screen 2
Diagnostic mode

Date/Time
The Date/Time function lets you verify date and time settings.

1. From the System Settings screen, select Date/Time to display the Set Date and Time screen (Figure E-5).

![Set Date and Time screen](image)

Figure E-5: Set Date and Time screen

2. Adjust the date and time with the + and - buttons; then Apply.
Pressure Units
The Pressure Units function lets you select the unit of measure for pressure display.

1. From the System Settings screen, select Pressure Units to display the Set Pressure Units screen (Figure E-6).

![Figure E-6: Set Pressure Units screen](image)

2. The active pressure unit is shown in white type. Select the desired pressure unit. The change is effective after you restart the ventilator.
Diagnostic mode

Restore Default Settings

The Restore Default Settings function lets you return ventilator settings to factory defaults. The factory defaults are listed in Chapter 11.

1. From the System Settings screen, select Restore Default Settings to display the Restore Default Settings screen (Figure E-7).

![Figure E-7: Restore Default Settings screen](image)

2. Select Restore Defaults.
Software Options
With the Software Options function, you enable a software option using a unique code specific to the option and the ventilator serial number. Options can also be enabled through the Respi-Link remote service program.

NOTE: Before installing an option, verify that the ventilator serial number matches the serial number shown in the Vent Info window (“Vent Info (ventilator information)” on page 6-18. If the serial numbers do not match, contact Philips.

1. From the System Settings screen, select Software Options to display the Enable Software Options screen (Figure E-8).

![Figure E-8: Enable Software Options screen](image)

2. Use the onscreen keypad to enter the code; then select Enter. The screen displays Enabled: followed by the name of the software option.

3. Repeat as needed to enable additional options.

4. Verify that the options are enabled by selecting Back to System Settings, then Back to Diagnostics Menu, then Service. The Vent Info window should show the new options.

5. Attach the option label as shown in Figure 3-5 on page 3-9.
Baud Rate
The Baud Rate function lets you set the baud rate for serial communications.

1. From the System Settings screen, select Baud Rate to display the Set Baud Rate for Serial Communications screen (Figure E-9).

![Set Baud Rate for Serial Communications screen](image)

Figure E-9: Set Baud Rate for Serial Communications screen

2. The active baud rate is shown in white type. Select the desired baud rate.
**Alarm Volume Escalation**

The Alarm Volume Escalation function lets you enable or disable volume escalation. When alarm volume escalation is Enabled and a high priority alarm is not responded to within 40 seconds, the ventilator alarm volume increases to maximum over an 20-second period.

When the Alarm Volume Escalation function is active and a touchscreen or button press is detected, the ventilator automatically returns the alarm volume to the user setting.

1. From the System Settings screen, select Alarm Volume Escalation to display the Set Alarm Volume Escalation screen (Figure E-10).

![Figure E-10: Set Alarm Volume Escalation](image)

2. The current setting is shown in the Information box at the top of the screen. If Alarm Volume Escalation is currently Disabled, you will see a selectable Enable button. If Alarm Volume Escalation is currently Enabled, you will see a selectable Disable button. Press the button to change the setting.

3. The new setting is applied after the V60/V60 Plus Ventilator is shut down and powered on again.

---

1. Available in Revision 2.30 software and above.
Diagnostic mode

Service

NOTE: All ventilator mode and alarm settings, alarm messages and significant events are automatically logged and retained, even when power is lost.

The Service screen lets you view the event log. Other service functions are for use by authorized service personnel.

Significant Event Log
The Significant Event Log contains data about clinically relevant ventilator occurrences, including alarms and setting changes. The time, date, and an identifier for event classification are included. A maximum of 2,000 event records are retained. The oldest events are overwritten first to allow recording of new events.

1. From the Service screen, select the Misc tab.

2. The Miscellaneous screen opens (Figure E-11). Select Significant Event Log.

Figure E-11: Miscellaneous screen
Diagnostic mode

3. The Significant Event Log opens (Figure E-12). Use the buttons on right side to navigate through the log.

Figure E-12: Significant Event Log screen
Calibrate the touchscreen X and Y coordinates as follows:

1. From the Diagnostics Menu, select Touch Screen Calibration. The Touch Screen Calibration screen is displayed (Figure E-13).

NOTE: If the Touch Screen Calibration button does not respond, press the Accept button on the top-right front of the ventilator to begin.

2. Follow the steps shown. Press on the middle of each target with a blunt, narrow object.

Figure E-13: Calibrate Touch Screen screen
Diagnostic mode

3. When prompted, touch the screen to exit calibration.
   If the calibration is not successful, have the ventilator serviced.

Exiting the diagnostic mode

Exit the diagnostic mode by turning off ventilator power with the ON/Shutdown key.
Diagnostic mode
Glossary

Ampere, a unit of current.

AC Alternating current.

Alarm Silence button Silences alarm sound for 2 minutes.

Alarm Volume escalation When enabled, this function becomes active if there is no response to a high priority alarm within 40 seconds. Ventilator alarm volume then increases to its maximum over a 20-second period.

Auto-Trak+ An optional feature that allows adjustments to trigger and cycle thresholds beyond Auto-Trak Sensitivity settings.

Auto-Trak Sensitivity A Respironics innovation in triggering and cycling that utilizes several different methods to provide enhanced sensitivity in the presence of leaks and changing breathing patterns.

AVAPS Average volume-assured pressure support. A ventilation mode in which pressure support is automatically adjusted to maintain the user-defined target tidal volume.

AVAPS Maximum IPAP Pressure See Max P.

AVAPS Minimum IPAP Pressure See Min P.

AVAPS Target Tidal Volume See V_T.

Average volume-assured pressure support See AVAPS.

Baseline As in baseline pressure. The pressure at end exhalation.

BPM Breaths per minute.

BTPS Body temperature (98 °F, ambient pressure), 100% saturated (with water vapor).

C-Flex A setting in CPAP mode, which enhances traditional CPAP by reducing the pressure at the start of exhalation.

cmH₂O Centimeters of water, a unit of pressure measurement.

Continuous positive airway pressure See CPAP.
Glossary

**CPAP** Continuous positive airway pressure. A ventilation mode that provides a single, continuous level of positive pressure to the patient and a control setting in that mode.

**Cycle** To end inspiration.

**dB(A)** Decibel, a unit of acoustic power.

**DISS** Diameter index safety standard, a standard for high-pressure gas inlet fittings.

**E-Cycle (Expiratory Cycle Sensitivity)** A control setting in Auto-Trak+. It determines the threshold at which the ventilator will transition from inspiration to exhalation.

**Elast.** See Elastance.

**Elastance** The elastic opposition to ventilation or the tendency of the lungs to resist inflation (elastance is the reciprocal of compliance).

**EPAP** Expiratory positive airway pressure. A control setting. The application and maintenance of pressure above atmospheric at the airway throughout the exhalation phase of positive-pressure mechanical ventilation.

**Estimated exhaled tidal volume** See $V_T$.

**Estimated minute ventilation** See $V_E$.

**Estimated patient leak** See Pt. Leak

**Estimated total leak** See Tot. Leak.

**ET** Endotracheal.

**Exhalation Port test** Performed to assess the leak flow rate through the exhalation port.

**Expiratory Cycle** See E-Cycle.

**Expiratory positive airway pressure** See EPAP.

**Flow** Flow rate, a setting in high flow therapy

**HFT** High flow therapy, a feature that provides a constant flow of mixed air and oxygen.

**HIP** High Inspiratory Pressure Alarm, an alarm setting.

**Hi Rate** High Rate Alarm, an alarm setting.

**Hi $V_T$** High Tidal Volume Alarm, an alarm setting.
Glossary

**hPa** Hectopascal, a unit of pressure measurement. 1 hPa is equal to 1 mbar, which is approximately equal to 1 cmH₂O.

**ID** Inner diameter.

**IEC** International Electrotechnical Commission.

**I:E ratio** Ratio of inspiratory to expiratory time.

**Inop** Inoperative.

**Inspiration:exhalation ratio** See I:E ratio.

**Inspiratory positive airway pressure** See IPAP.

**Inspiratory time** See I-Time.

**Inspiratory duty cycle** See \( \frac{T_i}{T_{TOT}} \).

**Intentional leakage** “Known,” quantifiable leakage that is a function of the mask.

**IPAP** Inspiratory positive airway pressure. A control setting. The application and maintenance of pressure above atmospheric at the airway throughout the inspiration phase of positive-pressure mechanical ventilation.

**ISO** International Organization for Standardization, a worldwide federation of national standards bodies.

**I-Time** Inspiratory time. The duration of inspiration during mechanical ventilation.

**L** Liter.

**LCD** Liquid crystal display.

**LED** Light-emitting diode.

**Limit** To prevent from exceeding a specified maximum value during a breath.

**LIP** Low Inspiratory Pressure Alarm, an alarm setting.

**Lo Rate** Low Rate Alarm, an alarm setting.

**Lo \( \dot{V}_E \)** Low Minute Ventilation Alarm, an alarm setting.

**Lo \( V_T \)** Low Tidal Volume Alarm, an alarm setting.

**Mandatory breath** A breath for which either the timing or volume is controlled by the ventilator. That is, the machine triggers and/or cycles the breath.

**Max E** Maximum elastance (volume assist). A control setting in PPV.
**Glossary**

**Max P** AVAPS Maximum IPAP Pressure. A control setting in AVAPS.

**Max P** Maximum Pressure. See PPV Maximum Pressure Limit.

**Max R** Maximum resistance (flow assist). A control setting in PPV.

**Max V** Maximum Volume. See PPV Maximum Volume Limit.

**Min P** AVAPS Minimum IPAP Pressure. A control setting in AVAPS.

**mL** Milliliter.

**mm** Millimeter.

**NIST** Non-Interchangeable Screw-Threaded. A connector for high-pressure gas inlet fittings.

**Noninvasive** Pertaining to a diagnostic or therapeutic technique that does not require the skin to be broken or a cavity or organ of the body to be entered. Mechanical ventilation via mask, nasal prongs, or mouthpiece.

**O₂** Oxygen (concentration). A control setting.

**OD** Outer diameter.

**PCV** Pressure-controlled ventilation. A ventilation mode that provides mandatory and spontaneous breaths with a set frequency, pressure, and inspiratory time.

**Peak inspiratory pressure** See PIP.

**Percentage of patient-triggered breaths** See Pt. Trig.

**PIP** Peak inspiratory pressure. The peak pressure for the previous inspiration.

**PPV %** A control setting in PPV. The percent of proportional pressure ventilation supplied by the ventilator.

**PPV** proportional pressure ventilation. A ventilation mode that delivers a pressure-controlled breath in proportion to the patient’s effort. The ventilator responds to patient instantaneous efforts, allowing the patient to determine when to start and end a breath, and how flow and pressure change as the patient breathes spontaneously.

**PPV Maximum Pressure Limit (Max P)** A control setting in PPV.

**PPV Maximum Volume Limit (Max V)** A control setting in PPV.

**Pressure-controlled ventilation** See PCV.

**Pressure-supported breath** A patient-triggered, pressure-targeted breath.
psi Pounds per square inch.

psig Pounds per square inch gauge (above atmospheric pressure).

Proportional pressure ventilation see PPV.

Pt. Leak The leak resulting from leaks around the mask or from unintentional leaks in the circuit. A monitored parameter shown when the intentional leak is known.

Pt. Trig Percentage of patient-triggered breaths. Patient-initiated breaths as a percentage of total breaths during the last 15 minutes.

Ramp Can be used to allow the patient to become accustomed to respiratory ventilatory therapy over time. Ramp will allow the pressure to linearly increase over a user-set period.

Rate (Respiratory Rate) Respiratory frequency, a control setting and monitored parameter.

Resist. See Resistance

Resistance The pressure drop across a pneumatic device (i.e., bacteria filter, patient circuit tubing) for a unit of flow when the volume of the device remains constant, i.e., cmH₂O/mL/sec.

Respiratory Rate (Rate) Respiratory frequency, a control setting.

Rise Time (Rise) The time required for a pressure-supported or pressure-controlled breath to reach its target pressure, a control setting.

RS-232 Serial data communications protocol.

SIS Sleeve Indexed System (Australia). A connector for high-pressure gas inlet fittings.

Spont indicator Denotes patient-initiated breathing.

Spontaneous breath A breath for which both the timing and volume are controlled by the patient. That is, the patient both triggers and cycles the breath.

Spontaneous/timed mode See S/T mode.

S/T mode Spontaneous/timed mode. A pressure support ventilation mode that ensures patients receive a minimum number of breaths per minute if their spontaneous breathing rate drops below the respiratory rate setting.

StandbySuspends ventilation and retains current settings when the clinician wants to temporarily disconnect the patient from the ventilator.
Glossary

Time Trigger Initiation of inspiration by the ventilator according to the Respiratory Rate setting.

Timed indicator Denotes machine-triggered (mandatory) breathing.

$T_i/T_{TOT}$ Inspiratory duty cycle. Inspiratory time divided by total cycle time, averaged over 8 breaths, a monitored parameter.

Tot.Leak Estimated total leak, both intentional and unintentional. A monitored parameter shown when the mask leak and type of exhalation port are not known.

Trigger To begin inspiration.

Trigger Sensitivity, a control setting in Auto-Trak+.

Trigger Sensitivity See Trigger.

Unintentional leakage Unpredictable leakage that cannot be quantified.

$V$ Volt, a unit of electrical potential or volume.

$\dot{V}$ Flow.

$\dot{V}_E$ Estimated minute ventilation. The product of tidal volume (spontaneous and timed) and rate (spontaneous and timed), a monitored parameter.

$V_T$ Estimated exhaled tidal volume, a monitored parameter and AVAPS Target Tidal Volume, a control setting in AVAPS mode.
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