Attention

Beijing Aeonmed Co., Ltd. (hereinafter referred to as "our company") holds the copyrights to this non-public published manual, and reserves the rights to keep it as a secure document. Refer to this manual when operating, maintaining and repairing products only. Only our company may provide permission for the use of or copies of this document to others.

Proprietary materials protected by the copyright law are included in this manual. No section of it may be duplicated, copied, or translated into other languages without prior written approval from our company who reserves the copyright.

Everything written within this manual is considered to be correct, but it is not a substitute for the exercise of professional judgment. Our company is not legally responsible for any mistakes printed within and/or any damages caused by incorrect connection and operation of equipment. Our company does not supply privileges endowed by the patent law to any other parties. Our company is not legally responsible for the results caused by patent law breaking or any rights of the third party violating.

Any user must read this article before using the products of our company. This article exposes the operating steps that must be read carefully. Improper use might endanger equipment or persons. The company will not undertake the responsibility for the safety, reliability and performance if equipment is used improperly. The company will not offer complimentary service for misused equipment.

Our company has the right to revise any content in this manual without notice; and has no obligation to update either hardware or software of the equipment described herein to the user or owner.
Warning for use

Welcome to use our products!

In order to use this product correctly and effectively, please read these operating instructions carefully and completely before using the product for the first time.

When using the product, always proceed in accordance with the information provided in these operating instructions on the basis of fully understanding the information in this manual.

This product is only for intended use as described in these operating instructions.

Only specially trained service professionals are authorized to perform the connection and service of this product.

For any situation in the use process, please contact with us. We will provide you with warm service.

Product specifications are subject to change without notification.
<table>
<thead>
<tr>
<th><strong>Manufacturer(holder):</strong></th>
<th>Beijing Aeonmed Co., Ltd.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Manufacturer(holder) address:</strong></td>
<td>Room 405, Basement 1 to 4th Floor of 901 Unit, Building 9, No.26 Outer Ring West Road, Fengtai District, Beijing 100070, China</td>
</tr>
<tr>
<td><strong>Facility:</strong></td>
<td>Beijing Aeonmed Co., Ltd.</td>
</tr>
<tr>
<td><strong>Facility address:</strong></td>
<td>Room 405, Basement 1 to 4th Floor of 901 Unit, Building 9, No.26 Outer Ring West Road, Fengtai District, Beijing 100070, China</td>
</tr>
<tr>
<td><strong>Service:</strong></td>
<td>Beijing Aeonmed Co., Ltd.</td>
</tr>
<tr>
<td><strong>Service Address:</strong></td>
<td>No.10 Chaobai Street, Yingbin Road West, Yanjiao Development Zone, 065201, Langfang, Hebei, China</td>
</tr>
<tr>
<td><strong>Tel:</strong></td>
<td>+86-10-83681616</td>
</tr>
<tr>
<td><strong>Fax:</strong></td>
<td>+86-10-63718989</td>
</tr>
<tr>
<td><strong>Service Line:</strong></td>
<td>+86 800-810-8333</td>
</tr>
<tr>
<td><strong>Website:</strong></td>
<td><a href="http://www.aeonmed.com">http://www.aeonmed.com</a></td>
</tr>
<tr>
<td><strong>E-mail:</strong></td>
<td><a href="mailto:service@aeonmed.com">service@aeonmed.com</a></td>
</tr>
<tr>
<td><strong>European Representative:</strong></td>
<td>Shanghai International Holding Corp. GmbH (Europe)</td>
</tr>
<tr>
<td><strong>Address:</strong></td>
<td>Eiffestrasse 80, 20537 Hamburg Germany</td>
</tr>
<tr>
<td><strong>Wechat Official Account:</strong></td>
<td><img src="image" alt="QR Code" /></td>
</tr>
</tbody>
</table>
# Table of Contents

1 **Introduction** ........................................................................................................ 1-1
   1.1 Manufacturer’s Responsibility ........................................................................ 1-1
   1.2 Operator’s Responsibility for Patient Safety .................................................. 1-1
   1.3 Definitions .......................................................................................................... 1-2
   1.4 Warnings, Cautions and Notes ........................................................................ 1-2
      1.4.1 Warnings ...................................................................................................... 1-2
      1.4.2 Cautions ....................................................................................................... 1-4
      1.4.3 Notes ............................................................................................................. 1-6
   1.5 Intended Use ........................................................................................................ 1-7
   1.6 Indications for Use ............................................................................................ 1-7
   1.7 Contraindication ............................................................................................... 1-8
   1.8 Abbreviations and Definitions ......................................................................... 1-8
   1.9 Frequently Used functions ............................................................................... 1-10
   1.10 Symbols ............................................................................................................. 1-10
   1.11 VG70 Ventilator Quick Start Guide ................................................................. 1-12

2 **System Overview** .................................................................................................. 2-1
   2.1 Ventilator Components .................................................................................... 2-1
   2.2 User Interface Components ............................................................................. 2-2
      2.2.1 GUI Screen Front Panel .............................................................................. 2-2
      2.2.2 GUI Screen Side Panel ................................................................................ 2-3
   2.3 Main Control Unit ............................................................................................ 2-4
      2.3.1 Front Panel ................................................................................................... 2-4
      2.3.2 Rear Panel .................................................................................................. 2-5
   2.4 Cart ..................................................................................................................... 2-6
   2.5 Humidifier ......................................................................................................... 2-6
   2.6 Cylinder Kit ....................................................................................................... 2-6

3 **CO₂ Module** .......................................................................................................... 3-1
   3.1 CO₂ Module Intended Use ................................................................................ 3-1
   3.2 CO₂ Module Specifications .............................................................................. 3-1
      3.2.1 General ......................................................................................................... 3-1
   3.3 System Assembly Instruction ............................................................................ 3-5
      3.3.1 Set-up .......................................................................................................... 3-5
      3.3.2 Placement of IRMA Probe .......................................................................... 3-7
   3.4 Pre-use Check .................................................................................................... 3-7
      3.4.1 Zeroing Procedure ..................................................................................... 3-7
   3.5 Alarms ................................................................................................................. 3-8
   3.6 Cleaning .............................................................................................................. 3-8
   3.7 Warnings ............................................................................................................ 3-8
   3.8 Cautions ............................................................................................................. 3-10
   3.9 Maintenance Information ............................................................................... 3-10

4 **Setup** .................................................................................................................... 4-1
   4.1 Connect Power Supply ..................................................................................... 4-1
1 Introduction

Review all information in this manual thoroughly before attempting to use the equipment.

This equipment must be used under the supervision of a physician.

1.1 Manufacturer’s Responsibility

Our company is responsible for the security; reliability and functions of the equipment, only when the following requirements are strictly adhered to:

- Only individuals authorized by our company may perform connection, adjustments and repairs.
- Necessary electrical equipment and the working environment must be in accordance with the national standards, professional standards and the requirements listed in this manual.
- Equipment must be used as instructed in this manual.

⚠️ CAUTION: This equipment is not for home use.

Our company will supply service information to help the customer, under the guidance of qualified technicians, to repair the equipment.

1.2 Operator’s Responsibility for Patient Safety

The operator of this ventilator must recognize their full responsibility for choosing appropriate ventilation settings to ensure proper ventilation and patient safety. The responsibility for the selection of the appropriate level of patient monitoring depends solely on the equipment operator.

All the monitoring information is for reference only; it should not be used as the sole basis for therapeutic or diagnostic decisions.

Whenever a patient is connected to the ventilator, constant attention by qualified medical personnel is required in order to provide immediate corrective action in case of a malfunction and/or alarm occurrence.

The company will provide functional block diagram at the user’s request for charge, accompanied by explanation on calibration method and other information, so as to help users let appropriate technical staff repair the equipment part that is allowed to be maintained by user as stipulated.
1.3 Definitions

This manual uses three special indicators to convey information of a specific nature. They include:

<table>
<thead>
<tr>
<th>Indicator</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="https://via.placeholder.com/15" alt="WARNING" /></td>
<td>Indicates a condition that can endanger the patient or the ventilator operator.</td>
</tr>
<tr>
<td><img src="https://via.placeholder.com/15" alt="CAUTION" /></td>
<td>Indicates a condition that can damage the equipment.</td>
</tr>
<tr>
<td><img src="https://via.placeholder.com/15" alt="NOTE" /></td>
<td>Indicates points of particular emphasis that make operation of the ventilator more efficient or convenient.</td>
</tr>
</tbody>
</table>

1.4 Warnings, Cautions and Notes

1.4.1 Warnings

![WARNING](https://via.placeholder.com/15) **WARNING:** Do not use the system until you have read and understood this manual including:

- All connections of the system
- All warnings and cautions
- Operation procedure of each and every component of the system
- Test procedure of each and every component of the system

![WARNING](https://via.placeholder.com/15) **WARNING:** The Ventilator System is intended for use by authorized and trained medical personnel only.

![WARNING](https://via.placeholder.com/15) **WARNING:** The users must familiarize themselves with the operation and use of this machine prior to first clinical use with a patient.

![WARNING](https://via.placeholder.com/15) **WARNING:** To ensure proper servicing and avoid the possibility of physical injury, only qualified personnel should attempt to service or make authorized modifications to the ventilator.

![WARNING](https://via.placeholder.com/15) **WARNING:** An authorized service engineer must first connect the ventilator and run our company’s connection procedure, which includes calibration of various system components, before you connect a patient to the ventilator.
**WARNING:** If a fault is detected in the ventilator so that its life support functions are no longer assured: start ventilation using an independent ventilation device (resuscitation bag) without delay, if necessary with PEEP and/or increased inspiratory $O_2$ concentration.

**WARNING:** Before activating any part of the ventilator, be sure to check the equipment for proper operation and, if appropriate, run PUT (pre-use test) as described in this manual, see section 5.

**WARNING:** The ventilator is not intended to be a comprehensive monitoring device and does not activate alarms for all types of dangerous conditions for patients on life-support equipment.

**WARNING:** Patients on life-support equipment must be appropriately monitored by competent medical personnel and suitable monitoring devices at all times.

**WARNING:** An alternative source of ventilation, such as manual respiratory equipment, should always be available when using the ventilator.

**WARNING:** Do not connect inspiratory or expiratory circuits to the exhaust port.

**WARNING:** Ensure that inspiratory and expiratory circuits are connected to the correct port before operation of equipment.

**WARNING:** The expiratory gas pathway may become contaminated with body fluids or expired gases during normal use, and the inspiratory gas pathway may become contaminated during fault condition, such as occlusion, breath hoses disconnection.

**WARNING:** Disposable breathing hoses shall not be reused. Reuse of the single use hoses can cause cross infection.

**WARNING:** Assure that hoses used have the appropriate resistance and compliance to ensure proper therapy.

**WARNING:** Do not disconnect the cable between the Main Control Unit and the GUI screen while Ventilator is operating.

**WARNING:** The ventilator must not be connected to any anti-static or electrically conductive hoses, tubing or conduit.
WARNING: Adding attachments or other components or sub-assemblies to the ventilator breathing system can change the pressure gradient across the ventilator breathing system and that such changes to the ventilation breathing system can affect the ventilator performance.

WARNING: Expiratory module is heated; use caution to avoid burns.

WARNING: Use caution when handling flammable or fragile components.

WARNING: Do not place containers of liquids (such as humidifier water reservoirs) on top of or above ventilator. Liquids getting into the ventilator can cause equipment malfunction with the risk of patient injury.

WARNING: To avoid risk of electric shock, this equipment must only be connected to a supply mains with protective earth.

1.4.2 Cautions

CAUTION: The breathing circuit must not be connected whenever the powers up and whenever a pre-use test is performed.

CAUTION: If the system test fails, do not use the system. Attempt to troubleshoot and fix the failure. If you are unable to fix the device, ask an authorized service representative to repair the device.

CAUTION: Check the ventilator periodically as outlined in this manual; do not use if defective. Immediately replace parts that are broken, missing, obviously worn, distorted, or contaminated.

CAUTION: Do not put ventilator into service until the patient setup is complete.

CAUTION: Measurements can be affected by mobile and RF communications equipment.

CAUTION: Do not use oxygen hoses that are worn, frayed, or contaminated by combustible materials such as grease or oils. Textiles, oils, and other combustibles are easily ignited and burn with great intensity in air enriched with oxygen.
CAUTION: Follow your hospital infection control guidelines for handling infectious material. Our company recognizes that cleaning, sterilization, sanitation, and disinfection practices vary widely among health care institutions. It is not possible for our company to specify or require specific practices that will meet all needs, or to be responsible for the effectiveness of cleaning, sterilization, and other practices carried out in the patient care setting.

CAUTION: Equipment not suitable for use in the presence of a Flammable Anesthetic mixture with Air or with Oxygen or Nitrous Oxide.

CAUTION: To avoid an electrical shock hazard while servicing the ventilator, be sure to remove all power to the ventilator by disconnecting the power source and turning off all ventilator power switches.

CAUTION: To avoid a fire hazard, keep matches, lighted cigarettes, and all other sources of ignition (e.g., flammable anesthetics and/or heaters) away from the ventilator and oxygen hoses.

CAUTION: In case of fire or a burning odor, immediately disconnect the ventilator from the oxygen supply, facility power and backup power source.

CAUTION: During operation, do not block: Speaker Holes, Exhaust Port, Air Inlet or Cooling Fan.

CAUTION: Do not use the Ventilator in a MRI environment.

CAUTION: The ventilator shall not be used in a hyperbaric chamber.

CAUTION: The ventilator shall not be used with helium or mixtures with helium.

CAUTION: Tip over hazard; use care when moving ventilator mounted to cart as device could tip over leading to injury or damage of equipment.

CAUTION: Do not use sharp objects to make selections on the LCD touch screen or panel.

CAUTION: Do not connect a VGA or USB interface while the system is in service.
CAUTION: The Network interface connection is for authorized service only.

CAUTION: Batteries should be removed if equipment will not be in service for more than 6 months. See Section 8.5 for battery replacement guidance.

CAUTION: Do not immerse the oxygen sensor or the connector in any type of liquid.

CAUTION: When ventilator is exposed to conditions outside the specified operating environment, allow 24 hours in normal environment before using.

CAUTION: Storage environment: -20°C ~+60°C and ≤95%RH.

CAUTION: Operating environment: 5°C ~40°C and 5%RH~95%RH.

CAUTION: Do not connect items that are not specified as part of the system.

CAUTION: The auxiliary outlet is only for the recommended humidifier; do not connect to any other equipment or an additional multiple socket outlets.

CAUTION: When using a humidifier, user should frequently check the water trap and look for water in the hose. If water is found in the hose, this water should be removed. Also, it is important the water trap is positioned in a way such that it is lower than the patient tubes.

CAUTION: Connecting electrical equipment to auxiliary outlet effectively leads to creating a medical equipment system, and can result in a reduced level of safety, make sure the ME SYSTEM comply with requirements of IEC 60601-1:2005. The user who connects is responsible for the standard for the requirements applicable to the medical equipment system.

1.4.3 Notes

NOTE: The user of this product shall have sole responsibility for any ventilator malfunction due to operation or maintenance performed by anyone not trained by our company.

NOTE: Usage of a filter on the expiratory side will increase the resistance of the patient circuit.
1 Introduction

⚠️ NOTE: In non-invasive (NIV) ventilation, the exhaled volume of the patient can differ from the measured exhaled volume due to leaks around the mask.

⚠️ NOTE: Do not sterilize or immerse the Mainstream CO₂ Adapter in any fluids. See Section 3 for proper use of Mainstream CO₂ Adapter.

⚠️ NOTE: All parts of the ventilator system are suitable for use within the patient environment.

⚠️ NOTE: All gas volume, flow, and leakage specifications in this manual are expressed at STPD (standard temperature and pressure dry), except when specified with another condition.

1.5 Intended Use

The VG70 Ventilator System is intended to provide continuous ventilation treatment to patients and monitoring of patients with respiratory failure or respiratory insufficiency, requiring respiratory support.

1.6 Indications for Use

The VG70 Critical Care Ventilator is an electronically controlled, electronically powered machine, which is used in ICU for the critical care, in Respiratory Department or Emergency Department for the rescue and therapy of the patient with respiratory insufficiency, and in other departments for providing respiratory support for the patient.

The Critical Care Ventilator should only be used by:

• Professional health care providers;

• Technicians that have received training in the use of this system.

The Critical Care Ventilator is applicable for the patient weighing at least 3 kg (7 lbs.), who require the following types of ventilatory support: Positive Pressure Ventilation, delivered invasively (by ET or Tracheotomy tube) or non-invasively (by mask) via Assist/Control, SIMV, CPAP and other modes of ventilation.

The Critical Care Ventilator is intended for use in hospital and hospital-type facilities. It may be used during intra-hospital transport provided that electrical power is supplied.
1.7 Contraindication

⚠️ **WARNING:** The Ventilator is not intended for use in areas with risk of explosion. Do not operate the ventilator in the presence of flammable anesthetics.

⚠️ **WARNING:** The Ventilator is not designed for use in an MRI environment. Do not use the Ventilator near an MRI machine; injury or equipment damage could result.

⚠️ **WARNING:** To avoid risk of electric shock, this equipment must only be connected to a mains supply with protective earth. A “hospital grade” cord must be used and connected to a “hospital grade” electrical outlet.

⚠️ **NOTE:** See Section 1.4 (Warnings, Cautions and Notes) for all environmental warnings.

1.8 Abbreviations and Definitions

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>(S)</td>
<td>Means Set Value</td>
</tr>
<tr>
<td>(M)</td>
<td>Means Measured Value</td>
</tr>
<tr>
<td>CPAP</td>
<td>Continuous Positive Airway Pressure (S)</td>
</tr>
<tr>
<td>f</td>
<td>Breath rate (frequency) in bpm, i.e. ventilation times per minute (S)</td>
</tr>
<tr>
<td>f_{spont}</td>
<td>Patient’s spontaneous respiratory frequency (M)</td>
</tr>
<tr>
<td>f_{total}</td>
<td>Total breath rate, i.e. the sum of breath rate f and spontaneous breath rate f_{spont} (M)</td>
</tr>
<tr>
<td>O_{2}</td>
<td>Inspiratory O_{2} concentration (S &amp; M)</td>
</tr>
<tr>
<td>I : E</td>
<td>The ratio of Inspiration to Expiration (M)</td>
</tr>
<tr>
<td>MV</td>
<td>Expiratory minute volume (M)</td>
</tr>
<tr>
<td>MV_{spont}</td>
<td>Spontaneously breathed minute volume (M)</td>
</tr>
<tr>
<td>MV_{leak}</td>
<td>Leakage minute volume (M)</td>
</tr>
<tr>
<td>Paw</td>
<td>Patient airway pressure (M)</td>
</tr>
<tr>
<td>PEEP</td>
<td>Positive End-Expiratory Pressure, which can improve the patient’s oxygenation (S &amp; M)</td>
</tr>
<tr>
<td>Symbol</td>
<td>Description</td>
</tr>
<tr>
<td>--------</td>
<td>-------------</td>
</tr>
<tr>
<td>PEEPi</td>
<td>Intrinsic Positive End-Expiratory Pressure (M)</td>
</tr>
<tr>
<td>P_{\text{insp}}</td>
<td>Upper pressure level in PCV mode (S)</td>
</tr>
<tr>
<td>P_{\text{mean}}</td>
<td>Mean airway pressure. This value is updated at the end of the last respiratory cycle, hence, is a continuous average (M)</td>
</tr>
<tr>
<td>P_{\text{peak}}</td>
<td>Airway pressure peak value during one ventilatory cycle (M)</td>
</tr>
<tr>
<td>P_{\text{plat}}</td>
<td>End-inspiratory airway pressure (M)</td>
</tr>
<tr>
<td>P_{\text{min}}</td>
<td>Minimum airway pressure (M)</td>
</tr>
<tr>
<td>P_{\text{sens}}</td>
<td>Pressure sensitivity (S)</td>
</tr>
<tr>
<td>P_{\text{supp}}</td>
<td>Pressure support (S)</td>
</tr>
<tr>
<td>P_{\text{high}}</td>
<td>Upper pressure level in BIVENT and APRV (S)</td>
</tr>
<tr>
<td>P_{\text{low}}</td>
<td>Lower pressure level in BIVENT and APRV (S)</td>
</tr>
<tr>
<td>T_{\text{max}}</td>
<td>Maximum inspiratory time (S)</td>
</tr>
<tr>
<td>T_{\text{insp}}</td>
<td>Inspiratory Time (S)</td>
</tr>
<tr>
<td>T_{\text{pause}}</td>
<td>Inspiratory Pause Time, to increase the inspiratory time to improve the patient’s oxygenation (S)</td>
</tr>
<tr>
<td>\dot{V}_{\text{sens}}</td>
<td>Trigger by flow rate (S)</td>
</tr>
<tr>
<td>V_T</td>
<td>Tidal volume of mechanical ventilation (S)</td>
</tr>
<tr>
<td>V_{te}</td>
<td>Expiratory tidal volume (M)</td>
</tr>
<tr>
<td>V_{li}</td>
<td>Inspiratory tidal volume (M)</td>
</tr>
<tr>
<td>E_{\text{sens}}</td>
<td>Expiratory trigger sensitivity (S)</td>
</tr>
<tr>
<td>ETCO_2</td>
<td>End-expiratory CO_2 concentration (M)</td>
</tr>
<tr>
<td>WOB</td>
<td>Work of breathing (M)</td>
</tr>
<tr>
<td>T_c</td>
<td>Time constant (M)</td>
</tr>
<tr>
<td>Leak%</td>
<td>Leakage percentage (M)</td>
</tr>
<tr>
<td>C_{\text{dyn}}</td>
<td>Dynamic compliance (M)</td>
</tr>
<tr>
<td>C_{\text{static}}</td>
<td>Static compliance (M)</td>
</tr>
</tbody>
</table>
### 1.9 Frequently Used functions

1. Power On / Off Switch
2. Connect patient hoses and gas supply
3. Pre-Use Test
4. Settings
5. Start Ventilation/Standby
6. Monitoring data
7. Alarm, Event/Alarm log
8. Calibration
9. Cleaning and disinfection
10. Breathing Circuit Components
11. System interconnections for gas supply
12. Humidifier and system interconnections
13. Nebulizer and system interconnections

### 1.10 Symbols

Instead of illustrations, symbols may be utilized. Not all of these symbols may necessarily appear on the equipment or in this User manual. The symbols include:

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
<th>Protection Class</th>
</tr>
</thead>
<tbody>
<tr>
<td>☀️</td>
<td>On (Power)</td>
<td>☐</td>
</tr>
<tr>
<td>⚪️</td>
<td>Off (Power)</td>
<td>☐</td>
</tr>
<tr>
<td>👤</td>
<td>Follow operating instructions</td>
<td>⚠️</td>
</tr>
<tr>
<td>🌡️</td>
<td>Protective earth ground</td>
<td>⚠️</td>
</tr>
</tbody>
</table>

- ☐: Protection Class Type B
- ☐: Protection Class Type BF
- ⚠️: Warning & Caution
- ⚠️: Dangerous voltage
<table>
<thead>
<tr>
<th>Action</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>EQUIPOTENTIAL</td>
<td>connection</td>
</tr>
<tr>
<td>Loudspeaker</td>
<td></td>
</tr>
<tr>
<td>Lock</td>
<td></td>
</tr>
<tr>
<td>Manufacturer</td>
<td>Date of production</td>
</tr>
<tr>
<td>Unlock</td>
<td></td>
</tr>
<tr>
<td>Serial Number</td>
<td></td>
</tr>
<tr>
<td>Inspiratory hold</td>
<td></td>
</tr>
<tr>
<td>Expiratory hold</td>
<td></td>
</tr>
<tr>
<td>Nebulization</td>
<td></td>
</tr>
<tr>
<td>Manual inspiration</td>
<td></td>
</tr>
<tr>
<td>Intelligent increase of oxygen</td>
<td></td>
</tr>
<tr>
<td>Standby</td>
<td></td>
</tr>
<tr>
<td>Waveform freeze</td>
<td></td>
</tr>
<tr>
<td>AC power</td>
<td>Internal Battery</td>
</tr>
<tr>
<td>USB device</td>
<td>Refer to documentation</td>
</tr>
<tr>
<td>Prompt message</td>
<td>Already online</td>
</tr>
<tr>
<td>Flow trigger</td>
<td>Pressure trigger</td>
</tr>
<tr>
<td>Adult</td>
<td>Manual trigger</td>
</tr>
<tr>
<td>NIV modes</td>
<td>Child</td>
</tr>
<tr>
<td>Main Menu</td>
<td>Invasive modes</td>
</tr>
<tr>
<td>Neonate</td>
<td>Alarm Silence Key</td>
</tr>
</tbody>
</table>
1.11 VG70 Ventilator Quick Start Guide

Review all information in the Operator’s Manual before attempting to use this equipment.

1. Connect Power Supply

Connect to AC Power Source, DC Power Source or Utilize Battery

2. Connect Gas Source

3. Power on Ventilator

Switch Ventilator on by turning to:

4. Technical Test

Technical Test in Progress...
5. Pre-use Test

6. Select New Patient

7. Prepare Patient Circuit

8. Connect CO₂ Module (optional)

9. Connect IRMA Airway Adapter (optional)
10. Set Appropriate Ventilation Settings
2 System Overview

2.1 Ventilator Components

The Critical Care Ventilator System consists of two required Components: a Main Control Unit and a Graphical User Interface (GUI). Optional Components available for the Critical Care Ventilator system are: Cart, Battery Backup Assembly, Patient Circuit Positioning Arm, Patient Circuit Assemblies and CO\textsubscript{2} module, etc.

![Figure 2-1 VG70 profile](image)

<p>| | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Main Control Unit</td>
<td>2</td>
</tr>
<tr>
<td>4</td>
<td>Water Trap</td>
<td>5</td>
</tr>
<tr>
<td>7</td>
<td>Test Lung</td>
<td>8</td>
</tr>
<tr>
<td>10</td>
<td>Nebulizer Connector</td>
<td>11</td>
</tr>
</tbody>
</table>

The GUI controls ventilator settings: Settings can be selected and adjusted by using a finger on the screen and/or the encoder knob. The GUI verifies that all combinations of settings are obtainable and will notify the user of any setting limitations.
Breathing parameters are continuously measured by transducers and controlled by a feedback system in the Breathing Delivery Unit. The ventilator responds to a difference between the actual measured value of a parameter and the preset or calculated value by adjusting gas delivery to achieve the target value.

### 2.2 User Interface Components

#### 2.2.1 GUI Screen Front Panel

![GUI screen front panel](image)

1. Alarm Lamp
2. Touch Screen
3. Encoder Knob
4. Alarm Silence Hard Key

The Alarm silence hard key enables the user to silence alarms for 2 minutes. Pressing a location on the GUI screen will either bring up a sub-menu or will highlight a ventilator parameter or shortcut key. Rotating the encoder knob when the ventilator parameter is selected allows the user to scroll through the available range.
2.2.2 GUI Screen Side Panel

![GUI screen side panel](image)

1. **Main cable**: This cable connects the GUI screen and the main control unit.

   ![CAUTION:](image) Do not disconnect this cable when the ventilator is operating.

2. **VGA interface**: For connection of a VGA display which will display the same information as the GUI screen. Display resolution of 1024*768, 60Hz is suggested.

   ![CAUTION:](image) Do not connect VGA interface when the system is in use on a patient.

3. **USB interface**: Only external storage devices of low power consumption are permissible for this USB interface.

   ![CAUTION:](image) Do not connect to the USB interface when the system is in service.

4. **Network Interface**: For connection to external equipment such as electronic health record. Refer to section 11.12, Communication/interface for additional information.

   ![CAUTION:](image) Do not connect Network Interface; only the qualified personnel can use it.
2.3 Main Control Unit

The main control unit is responsible for control of the ventilator. It has interfaces on the front and rear panels including pneumatic and electronic interfaces.

2.3.1 Front Panel

- **Expiratory module**: To remove the Expiratory module: Press the latch (5) on the right part of front cover (i.e. unlock it) and then take out the Expiratory module. After cleaning or high level disinfection, insert the module in the proper position until the locking latch returns to the locked state.

  **NOTE**: Use caution when inserting the expiratory module to avoid leakage. A system leak test must be done before the machine is put into patient use.

  **CAUTION**: Expiratory module is heated to prevent water condensation, use caution due to high temperature.

- **Exhaust port**: Patient expiratory gas is released through this port to room air.
2 System Overview

⚠️ **CAUTION:** Do not block this port.

⚠️ **CAUTION:** Do not connect inspiratory or expiratory limb circuit to this port.

3. Expiratory port

⚠️ **CAUTION:** Do not connect inspiratory limb circuit to this port.

4. Water trap cup: Collect condensed water to prevent it from going into expiratory valve.

5. Expiratory module latch: A latch that is used to lock or unlock the expiratory module.

⚠️ **WARNING:** Do not operate this latch or remove the Expiratory module while the Ventilator is in use on a patient.


7. Oxygen sensor cover

8. Inspiratory port: Delivers gas from the ventilator to the patient inspiratory limb hose.

⚠️ **CAUTION:** Do not connect expiratory limb circuit to this port.

### 2.3.2 Rear Panel

![Figure 2-5 Rear panel]
CAUTION: The Equipotential Terminal is used to connect various parts of the equipment or of a medical equipment system to the same potential. When connected, it shall comply with the IEC 60601-1.

WARNING: Never block the ports of the exhaust cooling fan and the air inlet! Clean the filter regularly! Ventilator should not be covered or positioned in such a way that that the operation or performance of the Ventilator is adversely affected (e.g. positioned next to a curtain that blocks the flow of cooling air, thereby causing the ventilator to overheat).

WARNING: Use caution when handling flammable and/or damageable components. Call for authorized service support when necessary.

CAUTION: CO₂ Module Connector is only for the specified CO₂ Module, and cannot be connected with other serial ports.

2.4 Cart

An optional cart may be used to mount the Ventilator. It applies to placing and moving the Ventilator. The cart includes an auxiliary AC panel.

2.5 Humidifier

An optional Humidifier, like the Fisher & Paykel MR850, or similar product, should be used with the Ventilator. Our company may supply humidifiers in your area – talk with our company’s Sales Representative if you need more information.

2.6 Cylinder Kit

An optional Gas Cylinder Mounting Kit is available that holds 2 US E cylinders (O₂).
3 CO₂ Module

3.1 CO₂ Module Intended Use

The mainstream CO₂ module is intended to be connected to the Ventilator for display of real time and derived monitoring data of CO₂.

The mainstream CO₂ module is intended to be connected to a patient breathing circuit for monitoring of inspired/expired gases during recovery and respiratory care. It may be used in the operating room, the intensive care unit, patient room and emergency medicine settings for adult and pediatric patients.

The CO₂ module is not intended to be used as the only means of monitoring a patient. It must always be used in combination with alternate monitoring systems.

3.2 CO₂ Module Specifications

3.2.1 General

<table>
<thead>
<tr>
<th>Description</th>
<th>Extremely compact infrared mainstream CO₂ probe.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dimensions (WxDxH)</td>
<td>38 x 37 x 34 mm (1.49” x 1.45” x 1.34”)</td>
</tr>
<tr>
<td>Cable Length</td>
<td>2.50 m ±0.02 m</td>
</tr>
<tr>
<td>Weight</td>
<td>&lt; 25 g (cable excluded)</td>
</tr>
<tr>
<td>Operating temperature</td>
<td>0 to 40°C / 32 to 104°F</td>
</tr>
<tr>
<td>Storage and transportation temperature</td>
<td>-40 to 75°C, -40 to 167°F</td>
</tr>
<tr>
<td>Operating humidity</td>
<td>10 to 95% RH, non-condensing</td>
</tr>
<tr>
<td>Storage and Transportation humidity</td>
<td>5 to 100% RH, condensing¹</td>
</tr>
<tr>
<td>Operating atmospheric pressure</td>
<td>525 to 1200 hPa (525 hPa corresponding to an altitude of 4572 m / 15000 feet)</td>
</tr>
<tr>
<td>Storage and transportation pressure</td>
<td>500 to 1200 hPa</td>
</tr>
<tr>
<td>Mechanical strength</td>
<td>Withstands repeated 1.8 m drops on a hard surface. Complies with requirements for road ambulances</td>
</tr>
</tbody>
</table>
Surface temperature (at ambient temp. 23°C) | Max 41°C / 106°F
--- | ---
Airway adapters | Disposable adult/pediatric: Adds less than 6 ml dead space; Pressure drop less than 0.3 cmH₂O @ 30 LPM. Disposable infant: Adds less than 1 ml dead space; Pressure drop less than 1.3 cmH₂O @ 10 LPM.

**NOTE 1:** After being in a condensing atmosphere, the unit shall be stored for more than 24 hours in an environment equivalent to the operating humidity. The humidity range 50 ~ 100% is valid within the temperature range of -40 to 40°C only.

### Gas analyzer

<table>
<thead>
<tr>
<th>Feature</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Probe</td>
<td>2-9 channel NDIR type gas analyzer measuring at 4 to 10 µm. Pressure, temperature and full spectral interference correction.</td>
</tr>
<tr>
<td>Calibration</td>
<td>Zeroing recommended when changing Airway adapter.</td>
</tr>
<tr>
<td>Warm-up time</td>
<td>Full accuracy within 10 seconds</td>
</tr>
<tr>
<td>Rise time (@ 10 l/min)</td>
<td>≤ 90 ms</td>
</tr>
<tr>
<td>Total system response time</td>
<td>&lt; 3 second</td>
</tr>
</tbody>
</table>

**Accuracy specifications – during standard conditions**

<table>
<thead>
<tr>
<th>Gas</th>
<th>CO₂ (%)</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO₂</td>
<td>0 – 15</td>
<td>±(0.2 vol% + 2% of reading)</td>
</tr>
<tr>
<td></td>
<td>15 – 25</td>
<td>Unspecified</td>
</tr>
</tbody>
</table>

**NOTE 1:** Gas concentration reported in units of volume percent.
Accuracy specifications – during all conditions 1)

<table>
<thead>
<tr>
<th>Gas</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO₂</td>
<td>±(0.3 vol% + 4% of reading)</td>
</tr>
</tbody>
</table>

⚠️ NOTE 1: The accuracy specification is valid for the operating temperature and humidity conditions specified, except for interference specified in the table “Interfering gas and vapor effects” below.

Interfering gas and vapor effects

<table>
<thead>
<tr>
<th>Gas or vapor</th>
<th>Gas level</th>
<th>CO₂</th>
</tr>
</thead>
<tbody>
<tr>
<td>N₂O 3)</td>
<td>60 vol%</td>
<td>- 1)</td>
</tr>
<tr>
<td>HAL 3)</td>
<td>4 vol%</td>
<td>- 1)</td>
</tr>
<tr>
<td>ENF, ISO, SEV 3)</td>
<td>5 vol%</td>
<td>+8% of reading 2)</td>
</tr>
<tr>
<td>DES 3)</td>
<td>15 vol%</td>
<td>+12% of reading 2)</td>
</tr>
<tr>
<td>Xe (Xenon) 3)</td>
<td>80 vol%</td>
<td>-10% of reading 2)</td>
</tr>
<tr>
<td>He (Helium) 3)</td>
<td>50 vol%</td>
<td>-6% of reading 2)</td>
</tr>
<tr>
<td>Metered dose inhaler propellants 3)</td>
<td>Not for use with metered dose inhaler propellants</td>
<td></td>
</tr>
<tr>
<td>C₂H₅OH (Ethanol) 3)</td>
<td>0.3 vol%</td>
<td>- 1)</td>
</tr>
<tr>
<td>C₃H₇OH (Isopropanol) 3)</td>
<td>0.5 vol%</td>
<td>- 1)</td>
</tr>
<tr>
<td>CH₃COCH₃ (Acetone) 3)</td>
<td>1 vol%</td>
<td>- 1)</td>
</tr>
<tr>
<td>CH₄ (Methane) 3)</td>
<td>3 vol%</td>
<td>- 1)</td>
</tr>
<tr>
<td>CO (Carbon monoxide) 4)</td>
<td>1 vol%</td>
<td>- 1)</td>
</tr>
<tr>
<td>NO (Nitrogen monoxide) 4)</td>
<td>0.02 vol%</td>
<td>- 1)</td>
</tr>
<tr>
<td>O₂ 4)</td>
<td>100 vol%</td>
<td>- 1)</td>
</tr>
</tbody>
</table>

⚠️ NOTE 1: Negligible interference, effect included in the specification “Accuracy, all conditions” above.

⚠️ NOTE 2: Interference at indicated gas level. For example, 50 vol% Helium typically decreases the CO₂ readings by 6%. This means that if measuring on a mixture containing 5.0
vol% CO₂ and 50 vol%. Helium, the measured CO₂ concentration will typically be \((1 - 0.06) \times 5.0\) vol% = 4.7 vol % CO₂.

⚠️ **NOTE 3:** According to the ISO 80601-2-55 standard.

⚠️ **NOTE 4:** In addition to the ISO 80601-2-55 standard.

⚠️ **CAUTION:** The presence of oxygen can cause some interference in the CO₂ measurement. This is known as spectral broadening, and must be compensated. The Ventilator performs the O₂ compensation automatically for IRMA CO₂. Use valid O₂ sensor, mount O₂ sensor and connect cable to ventilator correctly, maintain regularly. Otherwise etCO₂ accuracy may be affected.

⚠️ **CAUTION:** The presence of nitrous oxide can cause some interference in the CO₂ measurement. This is known as spectral broadening. The ventilator is not intended for use with nitrous oxide gas, and there is no compensation performed. Therefore, if nitrous oxide gas is used with the ventilator, the etCO₂ accuracy will be affected.
3.3 System Assembly Instruction

3.3.1 Set-up

1. Plug the IRMA connector into the CO₂ module connector on Ventilator rear panel and switch the power on.

2. Snap the IRMA probe on top of the IRMA airway adapter. It will click into place when properly seated.

3. A green LED indicates that the IRMA probe is ready for use.

4. Connect IRMA/airway adapter 15 mm male connector to the breathing circuit Y-piece.
5. Connect the IRMA/airway adapter 15 mm female connector to the patient’s endotracheal tube.

Alternatively, connect a HME (Heat Moisture Exchanger) between the patient’s endotracheal tube and the IRMA probe. Placing a HME in front of the IRMA probe protects the airway adapter from secretions and effects of water vapor and eliminates the need of changing the adapter. It allows free positioning of the IRMA probe as well.

6. Unless the IRMA probe is protected with a HME always position the IRMA probe with the status LED pointing upwards.
3.3.2 Placement of IRMA Probe

When connecting IRMA probe to an infant patient circuit it is important to avoid a direct contact between the IRMA probe and the infant’s body.

If, for whatever the reason, the IRMA probe is in direct contact with any parts of the infant’s body an insulation material shall be placed between the IRMA probe and the body.

⚠️ **WARNING:** The IRMA probe is not intended to be in patient contact.

3.4 Pre-use Check

Always verify gas readings and waveforms on the monitor before connecting the airway adapter to the patient circuit. Perform the tightness check of the patient circuit with the IRMA probe snapped on the IRMA airway adapter.

3.4.1 Zeroing Procedure

⚠️ **WARNING:** Incorrect probe Zeroing will result in false gas readings.

In order to secure high precision of the IRMA probe measurements the following zeroing recommendations should be followed.

Zeroing is performed by snapping a new IRMA airway adapter onto the IRMA probe without connecting the airway adapter to the patient circuit, and then using the host instrument to transmit a Zero reference command to the IRMA probe.

Special care should be taken to avoid breathing near the airway adapter before or during the Zeroing procedure. The presence of ambient air (21% O₂ and 0% CO₂) in the IRMA airway adapter is of crucial importance for a successful Zeroing. If an “ZERO_REQ” alarm should appear directly after a Zeroing procedure, the procedure must be repeated.
Always wait at least 10 seconds after changing the IRMA adapter before running the Pre-use test to allow the CO$_2$ sensor to warm up.

Zeroing needs to be performed only when an offset in gas values is observed, or when an unspecified accuracy message is displayed.

### 3.5 Alarms

The IRMA probe LED status is as follows:

<table>
<thead>
<tr>
<th>Light Status</th>
<th>Message</th>
</tr>
</thead>
<tbody>
<tr>
<td>Steady green light</td>
<td>System OK</td>
</tr>
<tr>
<td>Blinking green light</td>
<td>Zeroing in progress</td>
</tr>
<tr>
<td>Steady red light</td>
<td>Sensor error</td>
</tr>
<tr>
<td>Blinking red light</td>
<td>Check adapter</td>
</tr>
</tbody>
</table>

Refer to Section 8, Alarms and Troubleshooting, for EtCO2 low and high signal alarms.

### 3.6 Cleaning

The IRMA probe can be cleaned using a cloth moistened with maximum 70% ethanol or maximum 70% isopropyl alcohol.

Remove the disposable IRMA Airway Adapter prior to cleaning the IRMA probe.

⚠️ **CAUTION:** The airway adapters are non-sterile devices. Do not autoclave the devices as this will damage them.

⚠️ **CAUTION:** Never sterilize or immerse the IRMA probe in liquid.

### 3.7 Warnings

⚠️ **WARNING:** The IRMA probe is intended for use by authorized and trained medical personnel only.

⚠️ **WARNING:** The IRMA probe must not be used with flammable anesthetic agents.

⚠️ **WARNING:** Disposable IRMA airway adapters shall not be reused. Reuse of the single use adapter can cause cross infection.
**WARNING:** Used airway adapters shall be disposed of in accordance with local regulations for medical waste.

**WARNING:** Do not use the IRMA Adult/Pediatric airway adapter with infants as the adapter adds 6 mL dead space to the patient circuit.

**WARNING:** Do not use the IRMA Infant airway adapter with adults as this may cause excessive flow resistance.

**WARNING:** Measurements can be affected by mobile and RF communications equipment. Assure that the IRMA probe is used in the electromagnetic environment specified in this manual.

**WARNING:** Do not place the IRMA airway adapter between the endotracheal tube and an elbow as this may allow patient secretions to block the adapter windows and result in incorrect operation.

**WARNING:** To keep secretions and moisture from pooling on the windows sensor port, always position the IRMA probe in a vertical position with the LED pointing upwards.

**WARNING:** Do not use the IRMA airway adapter with metered dose inhalers or nebulized medications as this may affect the light transmission of the airway adapter windows.
3.8 Cautions

- **CAUTION:** Never sterilize or immerse the IRMA probe in liquid.

- **CAUTION:** The IRMA airway adapters are non-sterile devices. Do not autoclave the devices as this will damage them.

- **CAUTION:** Do not apply tension to the probe cable.

- **CAUTION:** Do not operate the IRMA probe outside the specified operating temperature environment.

3.9 Maintenance Information

The IRMA probe is permanently calibrated at the factory and requires verification in regular intervals using a service reference instrument.

- **WARNING:** Never put ventilator into service until patient setup is completed.

Clinical safety has been a major consideration in design of the machine, but operator should still be very cautious when operating the machine.

- **CAUTION:** In case any measured value seems suspect, operator should first examine the patient’s vital signs using other means, and then check the ventilator.
4 Setup

This section describes the connection and preparation of the ventilator.

4.1 Connect Power Supply

The machine can work with one of 2 power supply sources: internal battery and AC power supply. An icon on the right upper part of screen displays the supply being used. When operating on battery, all functions except the expiratory port heater, the cooling fan and the nebulizer are the same as under AC operation. The expiratory port heater, the cooling fan and the nebulizer are disabled in battery operation to improve battery run time. If “low battery” is displayed when the internal or extended battery is in use, the AC power supply must be connected to charge the battery, otherwise the Ventilator may lose power.

![Figure 4-1 Connect power supply cable to AC inlet](image)

After connecting the AC power supply, the AC power supply indicator will be shown indicating the battery is being charged. A typical charge period is 3.5 hours. The power supply indicator is lit yellow during periods of charging, and the light goes out when the battery is fully charged.

Remove AC Power supply cord from wall connection to disconnect ventilator from AC Mains.

4.2 Connect Gas Source

Hyperbaric Oxygen Inlet of the ventilator can be connected to multiple gas sources: bottled oxygen and central supply O₂. The gas source pressure must be between 280 ~ 600 kPa (41 ~ 87 psi). Low pneumatic pressure will impair some functions of the ventilator.
For Low-flow Oxygen Inlet of the ventilator, the gas source pressure must be less than 600kPa, and the flow is less than 15L/min.

There are diameter limits on the two inlets to prevent miss-connection.

Oxygen connected to the high pressure input ports of the ventilator will be used as Fresh Gas and will be supplied to the patient.

4.3 Connect Accessories

4.3.1 Connect Patient circuit

This figure shows the connection of patient circuit, including inspiratory port, water trap, Inspiratory tube, Y-piece, breathing hoses and expiratory filter.

⚠️ CAUTION: Assure patient hoses used have the appropriate resistance so that patient receives proper therapy.

⚠️ NOTE: When adding attachments or other components to the breathing system, the breathing resistance may increase, or the monitoring pressure of the patient connect port is higher than the actual pressure.
4.3.2 Connect Humidifier (optional)

Connect humidifier (1) into channel of mounting block (2).

4.3.3 Connect Patient Circuit Positioning Arm (optional)

Connect positioning arm (1) onto mounting block (2).
4.3.4 Connect User Interface Screen

Connect User Interface (1) to the location (2).

4.3.5 Connect Cylinder Kit (optional)

Connect O₂ cylinder (2) into the right side of holder (1). Secure cylinder using strap (3). Repeat for the other O₂ cylinder on the left side.

⚠️ **NOTE:** The kit includes only the mounting brackets and attachments. User must supply the Gas Cylinders with Gas Regulators and hoses compatible with the VG70 Gas Inlet Connections and at the proper Pressure per VG70 Specifications.
5 Pre-use Test

5.1 When to carry out pre-use test

- Before use of the Ventilator on a new patient
- After patient hose or patient filter replacement
- After maintenance or repair

⚠️ WARNING: Do not use the system until you have read and understood all the operation and maintenance manuals of the components.

⚠️ WARNING: If the system test fails, do not use the system. Attempt to troubleshoot and fix the failure. If you are unable to fix the device, ask an authorized service representative to repair the device.

⚠️ CAUTION: The following measures should be taken to minimize risks in the ventilator system.

5.2 Pre-use Test Procedure

After power on and technical test, the machine will enter the pre-use test following display of the power-on page. Items included: Gas supply test, Leak test, Flow sensor test, Pressure sensor test and Safety valve test in the power-on self-test: For details on testing, please refer to Section 6.1.

There is also a “Pre-Use Test” key on Standby screen. There are more test items in the “Pre-Use Test” page. Items included:

<table>
<thead>
<tr>
<th>Test Items</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Technical Test</td>
<td>After each system has completed its initialization technical tests will be performed, including: voltage checks at critical points in the circuitry; data collection necessary for system operation; test of communication between sub-systems; tests of measurement circuits and valve control circuits.</td>
</tr>
<tr>
<td>AC/Battery test</td>
<td>This test will verify whether the batteries can supply enough power to operate the ventilator normally. Please follow instructions as displayed.</td>
</tr>
<tr>
<td>Test Items</td>
<td>Remarks</td>
</tr>
<tr>
<td>-------------------------</td>
<td>-------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Gas supply test</td>
<td>Test will proceed when hyperbaric oxygen is functional.</td>
</tr>
<tr>
<td>Oxygen sensor test</td>
<td>This test requires that oxygen supply is available. If oxygen source is not available then a message “oxygen source is inadequate” and the test cannot be carried out.</td>
</tr>
<tr>
<td>Leak test</td>
<td>Internal leakage test.</td>
</tr>
<tr>
<td>Flow sensor test</td>
<td>Flow sensors function and accuracy test.</td>
</tr>
<tr>
<td>Pressure sensor test</td>
<td>Pressure sensors function and accuracy test.</td>
</tr>
<tr>
<td>Safety valve test</td>
<td>Safety valve function and accuracy test.</td>
</tr>
<tr>
<td>Patient circuit test</td>
<td>Circuit compliance value measurement.</td>
</tr>
<tr>
<td>CO₂ Sensor test</td>
<td>Performed if a CO₂ module is detected. An Alert will be posted if the test fails.</td>
</tr>
</tbody>
</table>

Below is a method for testing the function of the alarm system for conditions specified by IEC60601-2-12. Alarm system tests are to be performed at the user's discretion.

Perform the following procedure to verify operation of the Low MVe and High Airway pressure alarms:

1. Set the Power switch to ON.
2. Connect a breathing circuit and test lung to the ventilator.
3. Press Start Ventilation and ventilate with default settings except set O₂ to 21%.
4. After 5 breaths, observe the MVe reading on the display.
5. Set the MVe low alarm limit to a value greater than the observed MVe reading.
6. Verify that a low level MVe low alarm is present on the 3rd breath.
7. Return the MVe low alarm limit to original setting.
8. After 5 breaths, observe the Ppeak reading on the display.
9. Set the PAW upper alarm limit to a value lower than the observed Ppeak reading.
10. Verify that a low level High Airway Pressure alarm is present after 1 breath and that a high level High Airway Pressure alarm is present at the start of the 4th breath.
11. Return the PAW upper alarm limit to original setting.
12. Set the Power switch to OFF.
6 Ventilator Operation

⚠️ **WARNING:** Never put ventilator into service until patient setup is completed.

Clinical safety has been a major consideration in design of the machine, but operator should still be very cautious when operating the machine.

⚠️ **CAUTION:** In case any measured value seems suspect, operator should first examine the patient’s vital signs using other means, and then check the ventilator.

### 6.1 Starting Up

**Step 1: Connect to AC power supply**

Connect power supply cable to power supply socket on the wall, and the AC power supply indicator will be lit green.

![Connect power supply](image)

**Figure 6-1 Connect power supply**

**Step 2: Switching on**

To switch ventilator on, actuate the power switch of ventilator from "○" to "●". The machine will then be turned on; initialization of the GUI Display, Main Control unit and other systems will start, the power-on interface and then the company logo will be displayed.
Step 3: Technical test

After each system has completed its initialization, technical test will be performed, including voltages tests, data tests, communications tests, AD and DA converter tests, and valve control tests. The ventilator is not operational during this period.

The technical test should be performed for 10 seconds then wait for all results to be sent to GUI. If all tests pass, the technical test finished and continue to the pre-use test. If some test fail or waiting timed out, the failure information shall be displayed on screen, the “Test” and “Skip” button’s state will change to enable. User can choose “Test” to do technical test again, or choose “Skip” to skip technical test.
Step 4: Pre-use test

After the unit has finished the technical test, it shall enter into the pre-use test routine.

Test Items include: Gas supply test, Leak test, Flow sensor test, Pressure sensor test and Safety valve test. The pre-use test is an interactive test requiring the user to read and follow screen prompts. See Figure 6-4 for sample screen. Along the bottom of the screen shall be 2 buttons: Test and Skip.

Test Items shall be performed one by one. If all tests pass, it shall enter Standby screen. If some tests fail, the failed information shall be displayed on screen, the “Test” button’s state change to enable. Choose “Test” to do the test again, or choose “Skip” to enter Standby screen.
Figure 6-4 Pre-use Screen

⚠️ **NOTE:** The ventilator does not support patient ventilation during the pre-use, since the Inspiratory valve is closed, and Expiratory valve is opened.

### 6.2 Interface Layout

After the Power-on self-check is finished or the “Skip” key has been clicked. Press the blue key with the text “Start ventilation” ( ) to enter the currently selected ventilation mode and the blue key will change to orange with text “standby” on it ( ). After pressing and holding the standby key for 4 seconds, it will go back to “Start Ventilation” again.

#### 6.2.1 Standby Interface Layout

In the standby interface (see Figure 6-5), the user can set the information of the new patient, see the information of the previous patient, and do the pre-use test.
6.2.1.1 New Patient

a) Patient Settings

**Step 1:** Choose patient type (Adult or Child, default is adult)

**Step 2:** Ventilation type (Invasive or NIV, default is Invasive), Patient height (default is 150cm) and Ventilation mode. See example in Figure 6-6.

⚠️ **NOTE:** In the Ventilation Mode area, there is an "enter" key. Pressing the “Enter” key will enter the Mode of main menu.
b) **Patient Information**

Click the Patient Information key to enter patient information for a new patient. In this page, the user can enter the patient's name, Medical Record Number, Admission Date, Birth Date and Height (cm). There is a small keyboard on the right side of the Patient Information. The format of the admission date and birth date is YYYY/MM/DD, and the default date is the computer date. In the lower right corner, there is “Clear” key to clear all the information. See the example in Figure 6-7.

### Figure 6-6

![Patient Information Figure](image)

### Figure 6-7

![Patient Information Keyboard](image)

#### 6.2.1.2 Previous Patient

For the Previous Patient key, the same content displays as for a new patient, except the Patient Type and Vent Type appear on the Patient Settings page and the Patient information cannot be changed, as shown in Figure 6-8.
6.2.1.3 Pre-use Test

Click the Pre-Use test key to enter the pre-use test page. There are ten tests that must be done before the ventilator is connected to a patient: Technical test, AC/Battery Test, Gas Supply Test, Oxygen Test, Leak Test, Flow Sensor Test, Pressure Sensor Test, Safety Valve Test, Patient Circuit Test, and etCO2 Sensor Test. as shown in Figure 6-9.

There is a label with “√” at the front of each test item. Touch the test item can change the selected state of it. At the bottom of items there is a label with “√” and text “Select All”, touch it can select all test items.

⚠️ **NOTE:** Before the ventilator is connected to a patient, we suggest that the user do all tests.
6.2.1.4 Calibration

Click the calibration key to enter the calibration page, as shown in Figure 6-10. In this page, the user can calibrate $O_2$ sensor and flow sensor.

![Figure 6-10](image)

6.2.2 Ventilator Interface Layout

The ventilator interface can be divided into six parts: Parameter setup area, Short Cut keys area, Patient Measured Parameters area, Patient Waveforms area, Information area and User Message Prompts area, as shown in Figure 6-11.
Figure 6-11

Figure 6-12 shows the actual operational screen layout of the ventilator.
6.2.2.1 Information Area

The Information area includes seven sections: Ventilation Mode, Alarm Messages, Network and USB connection, Trigger, Patient Type and weight, AC and Battery indicators, and Time.

**Ventilation Mode area:** Displays the current mode of Ventilation.

**Alarm Messages area:** When there is no alarm message, this area is same background color as other screen areas; when a technical or functional alarm occurs, the background color will change to either red or yellow and text information will be displayed.

**Network and USB connection indicator area:** Displays the Network and USB connection status.

**AC and Battery indicators area:** Displays the AC, internal battery and extended battery connection status.

**Trigger area:** Displays the current ventilator trigger type. There are three trigger types: Pressure Trigger, Flow Trigger, and Manual Trigger. If there is currently no trigger in use, the trigger symbol will disappear.

⚠️ **CAUTION:** If the trigger sensitivity is set too high, a self-triggering (auto-triggering) condition may be reached. Triggering will then be initiated by the system and not by the patient. This should always be avoided by decreasing the trigger sensitivity. This is also important during transport as the movement of the body and the breathing system may lead to false triggering.

**Patient Type and weight area:** Displays the current patient’s type (adult or child) and current patient’s weight.

**Time area:** Displays the current time. There are two formats, 12-hour or 24-hour.
6.2.2.2 User Message Prompts Area

If no prompt message is displayed, this area will have the same background color as other parts of the screen. If a prompt message is available, it will have a flashing bulb icon in front of the relevant prompt message.

![Image of Standby screen](Figure 6-14)

6.2.2.3 Patient Waveform Area

At center of the screen, the Patient Waveforms area is the main display area. In the default state, this area will display three waveforms: Pressure waveform, Flow waveform and Volume waveform. See the example in Figure 6-15.

![Image of Patient Waveforms](Figure 6-15)
6.2.2.4 Patient Measured Parameters Area

This area displays parts of the monitored patient parameters which are very important. When in Standby mode, the monitoring values of all parameters will be displayed as “---”. The background color for Parameters with alarm limits will change between black and red when a high-level alarm occurs. The flash rate will be at 2 Hz ± 10 % and it will be synchronized with the high-level alarm displayed in the Alarm Message Area. All the parameters can be changed to other parameters by a sub menu which on the monitor. The parameters were separated into 3 groups, each group contain 2 parameters.

The parameters shall be divided into 3 groups: Group 1, 2 and 3. See example in Figure 6-16.

Figure 6-16

6.2.2.5 Ventilation Parameter Set-up Area

The ventilation parameter setup area is at the bottom of the screen. The breathing parameters settings necessary for the current ventilation mode are displayed in this area. See the example as shown in Figure 6-17. If the ventilation parameter setup items for the selected Ventilation mode do not fit in this space, the “More Settings” key will allow the user to access and change the other setup items.
6.2.2.6 Shortcut Keys Part

The ventilator has shortcut keys to access many ventilator operations, including Inspiratory Hold, Expiratory Hold, Nebulizer, Manual breath delivery, Suction, Print screen, Freeze, Screen Lock, Alarm Limits, Main Menu and Standby/Start Ventilation.
6.3 Operation of Main Manu

Click “Main Manu” of the shortcut key on the right side of the screen, the user can set Mode, Alarm Limits, Monitoring Data, Lung Mechanic, Log, System. Specific operation is as follows.

6.3.1 Ventilation Mode

6.3.1.1 Ventilation Mode Set-up

Step 1: Click in the shortcut keys area to enter the Main Menu, then click to enter mode Setup interface, as shown in Figure 6-19.
Step 2: Click the mode you want, for example:

Click [VCV] to enter the [VCV] mode setup page. The [VCV] key will become yellow as shown in Figure 6-20. The user can then set every parameter of VCV mode.

Step 3: Click [Accept], now VCV mode is set as the ventilation mode. See Figure 6-21.
NOTE: The setup procedures of other modes are similar to the one above.

WARNING: If “Accept” is not clicked, the screen will return to the main menu, and the last setup changes made will have no effect.

6.3.1.2 Mode Descriptions

Backup ventilation mode is included for: SPONT/CPAP+PSV, SIMV (VCV) +PSV, SIMV (PCV) +PSV, SIMV (PRVC) +PSV, and BIVENT+PSV. To setup the PSV mode, the user must set Psupp setting to a value not equal to 0, then “+PSV” will be added to the bottom of the mode name, as shown in Figure 6-22. Remember that mode changes are only in effect after Accept is pressed, as shown in Figure 6-23.
The spontaneous ventilation modes include: SPONT/CPAP + PSV, SIMV (VCV) + PSV, SIMV (PCV) + PSV, SIMV (PRVC) + PSV, and BIVENT + PSV. The backup ventilation mode will be PCV.

There are two measures for recovery from apnea: patient triggering and operator resetting.

When setting the above ventilation modes, the operator should set the backup ventilation mode. The default values and the range of the parameters are shown in the Table 6-1.

Table 6-1

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Factory default setup</th>
<th>Setup range</th>
<th>Adjustment step</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Adult</td>
<td>Child</td>
<td>Adult</td>
</tr>
<tr>
<td>Mode</td>
<td>PCV</td>
<td>PCV</td>
<td>V̇T (mL)</td>
</tr>
<tr>
<td>VT(mL)</td>
<td>400</td>
<td>80</td>
<td>50-2000</td>
</tr>
<tr>
<td>Parameter</td>
<td>Factory default setup</td>
<td>Setup range</td>
<td>Adjustment step</td>
</tr>
<tr>
<td>-----------------------------------</td>
<td>-----------------------</td>
<td>-------------</td>
<td>----------------</td>
</tr>
<tr>
<td></td>
<td>Adult</td>
<td>Child</td>
<td>Adult</td>
</tr>
<tr>
<td>f(bpm)</td>
<td>15</td>
<td>30</td>
<td>1-80</td>
</tr>
<tr>
<td>f(SIMV)(bpm)</td>
<td>10</td>
<td>20</td>
<td>1-40</td>
</tr>
<tr>
<td>f(NIV-T)(bpm)</td>
<td>15</td>
<td>30</td>
<td>4-20</td>
</tr>
<tr>
<td>PEEP/Plow(cmH₂O)</td>
<td>5</td>
<td>5</td>
<td>0-35</td>
</tr>
<tr>
<td>CPAP(cmH₂O)</td>
<td>5</td>
<td>5</td>
<td>2-20</td>
</tr>
<tr>
<td>O₂</td>
<td>40%</td>
<td>40%</td>
<td>21%-100%</td>
</tr>
<tr>
<td>P_{insp}(cmH₂O) in invasive modes</td>
<td>20</td>
<td>10</td>
<td>5-(70-PEEP)</td>
</tr>
<tr>
<td>P_{insp}(cmH₂O) in NIV-T</td>
<td>20</td>
<td>10</td>
<td>5-(50-PEEP)</td>
</tr>
<tr>
<td>T_p(s)</td>
<td>0</td>
<td>0</td>
<td>0-4</td>
</tr>
<tr>
<td>T_slope(s)</td>
<td>0.1</td>
<td>0.1</td>
<td>0-2</td>
</tr>
<tr>
<td>Trigger mode</td>
<td>V_sens</td>
<td>V_sens</td>
<td>V_sens,P_sens</td>
</tr>
<tr>
<td>P_{sens}(cmH₂O)</td>
<td>-3</td>
<td>-3</td>
<td>-20-0</td>
</tr>
<tr>
<td>V_sens(LPM)</td>
<td>2</td>
<td>2</td>
<td>0.5-20</td>
</tr>
<tr>
<td>T_{insp}(s)</td>
<td>1</td>
<td>0.6</td>
<td>0.2-9</td>
</tr>
<tr>
<td>I:E</td>
<td>1:2</td>
<td>1:2</td>
<td>1:10~4:1</td>
</tr>
<tr>
<td>P_{supp}(cmH₂O) in invasive modes</td>
<td>0</td>
<td>0</td>
<td>0-(70-PEEP)</td>
</tr>
<tr>
<td>P_{supp}(cmH₂O) in NIV-S/T</td>
<td>0</td>
<td>0</td>
<td>0-(50-PEEP)</td>
</tr>
<tr>
<td>P_{high}(cmH₂O)</td>
<td>15</td>
<td>15</td>
<td>5-60</td>
</tr>
<tr>
<td>P_{low}(cmH₂O)</td>
<td>5</td>
<td>5</td>
<td>0-35</td>
</tr>
<tr>
<td>T_{high}(s)</td>
<td>1</td>
<td>0.6</td>
<td>0.2-30</td>
</tr>
<tr>
<td>T_{low}(s)</td>
<td>3</td>
<td>1.4</td>
<td>0.2-30</td>
</tr>
<tr>
<td>E_sens</td>
<td>25%</td>
<td>25%</td>
<td>5%-80%</td>
</tr>
<tr>
<td>High Spont Insp Time</td>
<td>1.99 + (0.02x)</td>
<td>(1.99 + (0.02x)</td>
<td>0.4 sec to (1.99 + (0.02 x)</td>
</tr>
</tbody>
</table>
### Ventilator Operation

#### Parameter | Factory default setup | Setup range | Adjustment step
--- | --- | --- | ---
Adult | Child | Adult | Child
--- | --- | --- | ---
Patient height (cm) | 150 | 100 | 60-260 | 30-140 | 2
TC | OFF | OFF | ON, OFF | ON, OFF | --
Compliance compensation | ON | ON | ON, OFF | ON, OFF | --
Pipe diameter (mm) | 7.5 | 5.0 | 5-12 | 2.5-8 | 0.5

All modes in the mode menu include trigger type selection: pressure or flow. User can also set the value for $P_{sens}$ or $V_{sens}$ after choosing the trigger type. See Figure 6-24.

![Figure 6-24](image)

Tube compensation (TC) will be available only in pressure modes (PCV, SIMV (PCV), SPONT/CPAP and BIVENT). Select tube compensation “On” and the “On” key will become yellow as an acknowledgement, as shown in Figure 6-25.
Select the tube type in this interface: ET (Endotracheal Tube) or TT (Tracheotomy Tube), and the selected one will become yellow as shown in Figure 6-25.

Tube inside diameter (mm) and Tube compensation amount (%) may be modified after being clicked. Click “Accept” to store the new value.

When tube compensation is enabled, a message will be presented on the bottom of screen: TC ON, as shown in Figure 6-26. Also a tube icon will display in the information area, see Figure 6-27.

Tube inside diameter ranges are: 5 to 12mm for adult, 2 to 5-8mm for child; increment: 0.5mm for both adult and child.
Since the flow trigger is the default trigger mode, “pressure” is displayed on the trigger type menu. SIMV’s input interface is different from that of the other modes. The SIMV setup interface is shown in Figure 6-28, and the other modes in Figure 6-29.

![Figure 6-28](image)

After the “pressure” trigger mode is selected, the $P_{sens}$ key can be adjusted and the $V_{sens}$ key will become gray.

### 6.3.2 Alarm Limits

Click “Alarm Limits” in the “main menu” to enter the “Alarm Limits” menu interface, as shown in Figure 6-30.

![Figure 6-29](image)
The Alarm Limits setting includes:

- **Paw**: high and low limits of patient airway pressure (unit: cmH\(_2\)O)
- **MV\(_e\)**: high and low limits of minute volume (unit: LPM)
- **V\(_{te}\)**: low limit of tidal volume for Expiratory (unit: mL)
- **PEEP**: high and low limits of positive pressure at expiratory end (unit: cmH\(_2\)O)
- **Tapnea**: apnea duration (unit: second)
- **f\(_{spont}\)**: high limits of rate of spontaneous breaths (unit: breath per minute)
- **etCO\(_2\)**: high and low limits of CO\(_2\) concentration at the end of Expiration (unit: mmHg)

The limit setup ranges are shown in the following table:

<table>
<thead>
<tr>
<th>Limit Setting</th>
<th>Lower Limit</th>
<th>Upper Limit</th>
</tr>
</thead>
<tbody>
<tr>
<td>High limit of flow volume per minute</td>
<td>1 to 60L, OFF, 0.1 step, default 30L(_0)</td>
<td></td>
</tr>
<tr>
<td>Low limit of flow volume per minute</td>
<td>OFF, 0.1 to 40L, 0.1 step, default 1L for adult and 0.5L for child.</td>
<td></td>
</tr>
<tr>
<td>Low limit of tidal volume</td>
<td>5 to 400mL for child, 5 to 4000mL for adult. Step: 5mL for&lt;100mL (incl.), 10mL for &gt;100mL. Default: 250mL for adult and 50mL for child.</td>
<td></td>
</tr>
<tr>
<td>High limit of airway pressure</td>
<td>Not less than PEEP+5 or PEEP+PCV (PSV)(Phigh)+5; 5 to 80 cmH(_2)O, step 1 cmH(_2)O, default 40cmH(_2)O.</td>
<td></td>
</tr>
<tr>
<td>Low limit of airway pressure</td>
<td>OFF, 1 to 60 cmH(_2)O, step 1 cmH(_2)O, default 5 cmH(_2)O.</td>
<td></td>
</tr>
<tr>
<td>High limit of PEEP in Invasive Ventilation</td>
<td>1 to 35 cmH(_2)O, OFF, step 1 cmH(_2)O, default 10 cmH(_2)O.</td>
<td></td>
</tr>
<tr>
<td>Low limit of PEEP in Invasive Ventilation</td>
<td>OFF, 1 to 35 cmH(_2)O, step 1 cmH(_2)O, default OFF.</td>
<td></td>
</tr>
<tr>
<td>Parameter</td>
<td>Default Setting</td>
<td></td>
</tr>
<tr>
<td>---------------------------------------</td>
<td>-----------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>High limit of PEEP in NIV</td>
<td>1 to 20 cmH₂O, OFF, step 1 cmH₂O, default OFF.</td>
<td></td>
</tr>
<tr>
<td>Low limit of PEEP in NIV</td>
<td>OFF, 1 to 20 cmH₂O, step 1 cmH₂O, default OFF.</td>
<td></td>
</tr>
<tr>
<td>High limit of apnea duration</td>
<td>10 to 60s, OFF, step 1s, default 20s.</td>
<td></td>
</tr>
<tr>
<td>High limit of spontaneous breath rate</td>
<td>10 to 80 BPM, OFF. Default: OFF.</td>
<td></td>
</tr>
<tr>
<td>High limit of etCO₂</td>
<td>0.1%<del>13.3% (1mmHg</del>100mmHg or 0.1kPa~13.3kPa), default 6.5% (49mmHg or 6.5kPa).</td>
<td></td>
</tr>
<tr>
<td>Low limit of etCO₂</td>
<td>OFF, 0.1%<del>13.2% (OFF, 1mmHg</del>99mmHg or OFF, 0.1kPa~13.2kPa), default 4.0% (30mmHg or 4.0kPa).</td>
<td></td>
</tr>
</tbody>
</table>

⚠️ **CAUTION:** Do not set alarm limit parameter to extreme values that can render the alarm system useless.

⚠️ **NOTE:** The “default” values are manufacturer-configured alarm presets, user-configured alarm presets can be made different from the manufacturer-configured alarm presets in [Configurations] menu.

⚠️ **WARNING:** A potential hazard can exist if different ALARM PRESETS are used for the same or similar equipment in any single area, e.g. an intensive care unit or cardiac operating theatre.

⚠️ **WARNING:** When selecting “New patient”, confirm all alarm limits parameters are appropriate prior to use on each patient

⚠️ **NOTE:** To set alarm parameter to “OFF”, set the data display to the lowest or highest value. Press the encoder knob, turn encoder knob one more click and confirm by pressing encoder knob once more.

⚠️ **NOTE:** All alarm limit setting parameters are retained during power interruption and can be restored when power returns. Alarm volume can be adjusted to 5 levels: 20%, 40%, 60%, 80%, 100%. The default volume is 20%.

⚠️ **NOTE:** If [Alarm volume] requires adjustment, the procedure is as follows (the same procedure is used for other alarm parameter setup):

**Step 1:** Click “Alarm volume” when the key’s background color becomes yellow, then you can adjust the level of alarm volume, as shown in Figure 6-31.
Step 2: Turn the encoder knob left or right to adjust the alarm volume, press the encoder knob to confirm when proper value is reached. Background color will return to its original color, as shown in Figure 6-32.

⚠️ CAUTION: If the encoder knob is not pressed at the end, the system will go back to the original value after 10 seconds, i.e. the new setting will have no effect.

⚠️ CAUTION: In the case of an alarm during operation, the following cases may have occurred:

1) Improper breathing parameter setting or alarm limit setting;

2) Leakage in patient circuit; turn off the machine first and then check. In case of no resolution, contact service representative.

3) Problems with patient;

4) Power supply failure or ventilator failure.

In case there is not sufficient gas volume given to the patient, disconnect the ventilator from the patient, use artificial respiration or other emergency devices for the patient. Check the machine thoroughly.
6.3.3 Monitoring Data

Click “Monitoring data” on main menu to enter “Monitoring data” interface. The parameters are listed in four columns in this interface, as shown in Figure 6-33. All parameters will be also shown in the main interface.

![Figure 6-33](image)

6.3.4 Lung Mechanics

Click “Lung Mechanics” on “main menu” to enter “Lung Mechanics” page. On this page, you can select the test items on the left (from the top down): Rinsp, C static and PEEPi. The results of last test for the six parameters are listed on the right, as shown in Figure 6-34.
Step 1:

Click “Rinsp” to enter the test menu. The basic information is displayed in the middle of the screen, which includes the result of last and current inspiratory Resistance measurement. The result with date and time of last measurement is on the left, and the result of current measurement with time and date is on the right, as shown in Figure 6-35. If the measurement is not started, the results are “--”. 

![Figure 6-35](image-url)
Step 2:

When the measurement is completed, current measured values with date and time will be present. “Start” key appears again for another measurement.

⚠️ **NOTE:** The test procedures of parameter C static and PEEPi are similar to “Rinsp”.

### 6.3.5 Log

Click “Log” on “main menu” to enter log menu. There are two keys on the left of the page: Event/Alarm and trend, as shown in Figure 6-36.

![Figure 6-36](image)

**Step 1:**

Click “Event/Alarm” on log menu to enter the submenu, as shown in Figure 6-37.

![Figure 6-37](image)
The middle area of this page is the message area. It can store up to 1000 messages, including event messages and alarm messages. All messages will be listed in time sequence. The top is the latest event or alarm message, and the bottom is the oldest. Use the scroll bar to check all the messages. An asterisk (*) in front of an alarm message indicates that alarm message was not displayed in the alarm message area. As shown in Figure 6-38. The event/alarm log records all alarms and most actions.

<table>
<thead>
<tr>
<th>Date</th>
<th>Time</th>
<th>Event</th>
<th>Alarm</th>
</tr>
</thead>
<tbody>
<tr>
<td>2013-02-26</td>
<td>11:40</td>
<td>Communications Error 1005</td>
<td></td>
</tr>
<tr>
<td>2013-02-26</td>
<td>10:02</td>
<td>Communications Error 1005</td>
<td></td>
</tr>
<tr>
<td>2013-02-26</td>
<td>10:02</td>
<td>Communications Error 1006</td>
<td></td>
</tr>
<tr>
<td>* 2013-02-26</td>
<td>15:05</td>
<td>Apneia III</td>
<td></td>
</tr>
<tr>
<td>2013-02-25</td>
<td>15:04</td>
<td>Communications Error 1006</td>
<td></td>
</tr>
<tr>
<td>2013-02-25</td>
<td>15:04</td>
<td>Communications Error 1005</td>
<td></td>
</tr>
<tr>
<td>2013-02-25</td>
<td>11:40</td>
<td>Communications Error 1006</td>
<td></td>
</tr>
<tr>
<td>2013-02-25</td>
<td>11:40</td>
<td>Communications Error 1005</td>
<td></td>
</tr>
</tbody>
</table>

Figure 6-38

Below is the message area of the Settings area resulting from highlighting an alarm. All settings will be given here, as shown in Figure 6-39.

<table>
<thead>
<tr>
<th>Settings:</th>
<th>Patient:</th>
<th>Mode:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adult</td>
<td>1 bpm</td>
<td>15</td>
</tr>
<tr>
<td></td>
<td>O2 %</td>
<td>40</td>
</tr>
<tr>
<td></td>
<td>PEEP cmH2O</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>TpO2</td>
<td>1.0</td>
</tr>
<tr>
<td></td>
<td>VT mL</td>
<td>400</td>
</tr>
<tr>
<td></td>
<td>PEEP cmH2O</td>
<td>20</td>
</tr>
<tr>
<td>PCV</td>
<td>0.1</td>
<td>Frurog cmH2O</td>
</tr>
<tr>
<td></td>
<td>EEEEn %</td>
<td>25</td>
</tr>
<tr>
<td></td>
<td>HSMV bpm</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>CPAP cmH2O</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>PEEP cmH2O</td>
<td>15</td>
</tr>
</tbody>
</table>

Figure 6-39

⚠️ NOTE: Alarm/Event Log data is retained during a power interruption and can be viewed when power returns.

Step 2:

Click “Trend” to enter trend interface. There are two trend types: graphical trend and tabular trend, See Figure 6-40 & Figure 6-41. The default is graphical trend. Choose the tabular trend to get more information.

In “Graphical Trend”, the first page displayed is trend map of pressure-related parameters, including Ppeak, Pplat, PEEP, with timeline within 72h. See Figure 6-40.
Each graph shall have parameter as a title, the units of the parameter and range scale. Six time bases shall be 1, 3, 6, 12, 24, and 72. The displayed parameter for each Trend graph shall be selected from a pop-up menu that will give the user 8 parameters to choose from. The 8 parameter choices in the pop-up menu shall be configured in the Systems tab. “Zoom In” and “Zoom Out” keys shall be available so that there is better resolution of the Graphs. “Left Arrow” and “Right Arrow” keys shall be available to move a measuring line to the left or right. The measured value shall be displayed to the right of the parameter label on the right side of each graph. “Zoom In” shall decrease the time base. “Zoom Out” shall increase the time base. A Tabular trend key shall be available for the user to select.

**NOTE:** The X-axis is timeline for all trend maps and Y-axis is the corresponding parameter units.

The waveform on trend map is refreshed from left to right, with latest data on the very left. Y-axis is set for full scale range display.
6.3.6 System

Click “System” key to enter the System interface, as shown in Figure 6-42.

![Figure 6-42](image)

There are four keys on the left of the page: Settings, configurations, machine information and service.

6.3.6.1 Settings

Click "settings" to enter the settings interface, as shown in Figure 6-43.

![Figure 6-43](image)
You can set the Gas standard, time and date, unit setting, and compliance compensation in this page,

1. **Gas standard**

There are two Gas Standard options available to the user: BTPS (Body Temperature and Pressure Saturated) and ATP (Ambient Temperature and Pressure). When selected, the key background color shall be yellow. The default is BTPS and it is chosen with a dot.

2. **Compliance compensation**: ON or OFF.

3. **Dead space compensation**: ON or OFF.

4. **Unit setting**

Pressure unit: Two options: cmH₂O and hPa. When a unit is selected, all pressure units are converted to this unit, such as pressure unit in parameter setting area and parameter monitor area. Default: cmH₂O.

Weight unit: Two options: kg and lb. When a unit is selected, all body weight units are converted to this unit. Default: kg

CO₂ unit: Three options: mmHg, kPa and %. When a unit is selected, all CO₂ units are converted to this unit. Default: mmHg.

5. **Time and date**

Date format: YYYY/MM/DD; Time format: 24hours.

When the date/time format is modified, all date/time areas on the screen will be modified simultaneously. Modify date by clicking directly the numerical value at electronic calendar.

6.3.6.2 **Configurations**

Click “Configurations” key to enter the configuration interface, as shown in Figure 6-44.

There are three choices in this page: Graphic Trend, Screen Brightness and Site configuration.
6.3.6.2.1 Graphic Trend

Click “Graphic Trend” to the “Graphic Trend” interface, as shown in Figure 6-45.

On this page, the Patient Measured Parameters can be set. All graphic trends recorded are chosen from this screen. Trend 1-4 are the maps on the first page, and trend 5-8 are the maps on the second page. Below the 8 parameter, there is a monitoring parameter list, including 18 parameters. The Trend 1-8 can be changed according to the need from the parameter list. When a new parameter is chosen, the parameter on the main screen will be replaced at the same time.
6.3.6.2.2 Screen Brightness

Click “Screen Brightness” to enter this page, and click the screen to choose day or night. As shown in Figure 6-46. Day is the default.

6.3.6.2.3 Site Configuration

Step 1

Click “Site Configuration” to enter this page. Set a 4 digit numeric password in the dialog box to have protected access to set Configuration. As shown in Figure 6-47.
Step 2

After entering the password correctly, configure the sub menu: As shown in Figure 6-48.

1) Ventilation;
2) Alarms;
3) TC;
4) Monitoring;
5) Others;
6) Network;
7) Load/Save.

![Main Menu](image)

Figure 6-48

(1) Ventilation

a. Settings

Click “Ventilation” key, the background color will change to yellow and the default page “settings” will be displayed, as shown in Figure 6-48. User can configure this page for 1) Vent Type, 2) Ventilation Mode, 3) Patient Type, 4) Start-Up value for Vt based on weight?, and 5) Inspiratory time. Touching the key to choose the relative setting will remove the previously selected configuration. Only one choice shall be selected per setting. A selection shall also be provided to enable the user to restore the initial settings to factory defaults. Pressing the “Restore” key shall immediately restore all displayed choices to their factory default.

I. Vent type: Invasive or NIV (Non-Invasive), the factory default is Invasive.

II. Ventilation Mode: VCV, PCV, PRVC, SIMV (VCV), SIMV (PCV), SIMV (PRVC), BIVENT, and SPONT/CPAP for invasive mode, and for NIV are NIV/CPAP, NIV-T and NIV-S/T. The factory default is PCV for Invasive mode and NIV/CPAP for NIV mode.

III. Patient Type: Adult or Child. The factory default is Adult.

IV. Start-up value for Vt based on weight?: Yes or No. The factory default is No.
V. Height: Adult or Child. The default for adult is 150cm and for child is 100cm.

VI. Inspiratory time: Tinsp or I:E. The factory default is Tinsp.

After setting all the values needed, press the “Save” key, otherwise all the settings are lost. Another key “Restore” is under the “Save” key, pressing it restores all displayed choices to their factory default.

b. Parameters

Selecting “Parameters” key, will display a parameters configuration page. This includes the adult and child initial parameter settings. The user can configure for 1) Vt, 2) f, 3) Tinsp, 4) Pinsp, 5) Tslope, 6) PEEP, and 7) Psupp 8) O2. As shown in Figure 6-49.

![Main Menu](image)

Figure 6-49

Each setting key represents a location that the user can select. Touching the key will change the background color of the key, indicating that it has been chosen and can be changed by rotating the encoder knob left or right to adjust the value. When the selection reaches its maximum or minimum allowable setting, further rotation shall result in the minimum or maximum value displayed continuously. Pressing the key or the encoder knob again confirms the change.

⚠️ NOTE: Touching any other keys (or touchable area), will deselect the change and the parameter will be restored to the original setting. The background color will return to normal if the encoder knob is not pressed within 10 seconds

After setting all the values needed, the user will press the “Save” key. Pressing the “Restore” key restores all displayed settings to their factory default.

About the range of the parameters setting, please refer to Table 6-1.

(2) Alarms
Click the Alarms key to enter the alarms configuration page. There are two pages and six configurable alarms on this screen: 1) Paw, 2) etCO2, 3) f spont, 4) PEEP, 5) Tapnea, 6) MVe. See Figure 6-50.

![Figure 6-50](image1)

Each setting key represents a location that the user can select. Touching the key to choose or change the value, enter the upper and lower limits needed for each alarm, then press the Save key to confirm. There are three configurable alarms available for the user in the second page: 1) MVe, 2) Vte. You can change the alarm limits based on whether the patient is an adult or child, then press the Save key to save the new alarm limits.

Also there is a Restore key under the Save key. Pressing the Restore key restores all displayed settings to their factory default. As shown in Figure 6-51.

![Figure 6-51](image2)
(3) TC

Click the TC (Tube Compensation) key to enter the TC configuration page. In this page, you can configure for 1) Tube Compensation, 2) Tube Type, 3) Compensation %, 4) Diameter Adult and 5) Diameter Child, as shown in Figure 6-52.

![Main Menu](image)

**Figure 6-52**

1) Tube compensation: ON or OFF, the factory default is OFF.

2) Tube Type: ET or TT, the factory default is ET.

3) Compensation %, including Diameter Adult or Diameter Child. The range for Compensation is 0 to 100 in 1% increments and the factory default is 0. Diameter Adult range is 5.0 to 12.0 mm and the default is 7.5 mm. Diameter Child range is 2.5 to 8.0 mm and the default is 5.0 mm.

After finishing all the settings, the user shall press the Save key. Pressing the Restore key restores all displayed settings to their factory default.

(4) Monitoring

Click the Monitoring key to enter the Monitoring configuration page. On this page, you can configure whether the monitoring is ON or OFF, as shown in Figure 6-53.
(5) Others

Click the “Other” key to enter the others configuration page. On this page, you can choose the oxygen supply type, either HPO or LPO. The factory default shall be HPO. As shown in Figure 6-54.

When the Oxygen Supply Type is LPO, the ventilator should:

- Cancel the $O_2$ high/low concentration alarm and low Oxygen Supply Pressure alarm.
- $O_2$ parameter adjustment shall be disabled.
- The nebulizer, suction, $O_2$ sensor calibration functions shall be invalid.
- No Oxygen Supply Pressure monitor and calibration.
- When performing Pre Use Test, skip Gas supply test and Oxygen sensor test.
(6) Network

Ventilator data is shared by NetWork. As shown in Figure 6-55.

![Main Menu](image)

Figure 6-55

(7) Load/Save

Click the "Load/Save" key to enter the Load/Save configuration page. On this page, the load button shall be load the user configuration to the machine, and the save button shall be save the current configuration to the file which you saved. As shown in Figure 6-56.

![Main Menu](image)

Figure 6-56
6.3.6.3 Service

Click “Service” to enter the “Service” page. On this page, the user needs to enter the password, as shown in Figure 6-57.

Figure 6-57

Input the correct password to enter. There are six choices on this page: Calibration, Event/alarm log, Machine Information, Language, Test Page, Update and Optional. The default page is Calibration. See Figure 6-58.

Figure 6-58
6.3.6.3.1 Calibration

The calibration choices include: Pressure Sensor Calibration, Flow Sensor Calibration, O2 Sensor Calibration, CO2 Sensor Calibration, Inspiratory valve Calibration, Expiratory Valve Calibration, Atmospheric Sensor Calibration, Touch Screen Calibration, Leak Test and Breath Circuit test, as shown in Figure 6-59.

(1) Pressure Sensor Calibration: Click “Pressure Sensor” to enter the calibration interface. A message is displayed: “This step is to zero the pressure sensor. Please remove the breathing circuit from the ventilator before calibration.” A legend is displayed as well, as shown in Figure 6-59.

Click “Start” to start pressure sensor calibration. A progress bar and a message “Calibration in progress, please wait” will be displayed as shown in Figure 6-60. After calibration, the result will appear: Calibration succeeded or Calibration failed. If failed, restart the calibration.

NOTE: During this period no other operation can be performed. Clicking other areas will have no response.
(2) Flow Sensor Calibration: Click “Flow Sensor” to enter the interface. A message is displayed: “This step is to calibrate the flow sensor. Please connect the insp. Port and Exp. Port directly with a tube” as shown in Figure 6-61.

Click the “Start” button to start flow sensor calibration, the remaining procedure is the same as the pressure sensor calibration.

(3) O₂ Sensor Calibration: Click “O₂ Sensor” to enter the interface. A message displayed: “Please verify that the oxygen source are connected correctly. Verify that the gas inlet pressure is within specification.” A legend will also be shown. There are two keys below the legend: “Start 21%” and “Start 100%”. Choose the needed one and click, as shown in Figure 6-62. The remaining procedure is the same as described above.
(4) CO₂ sensor Calibration: Click “CO₂ Sensor” to enter the interface, as shown in Figure 6-63. A message displayed: “Disconnect the CO₂ sensor with the adapter from breathing circuit and ensure it is in ambient air. Wait 1 minute for warm up after the unit is powered on or after connecting an airway adapter. Press “Zero” when the State-Area turns green”. Please follow the prompt message to calibrate.

(5) Inspiratory Valve Calibration: Click “Inspiratory Valve” to enter the interface. A message displayed: “This step is to calibrate the inspiratory valve. Please connect the Insp. Port and Exp. Port directly with a tube”, as shown in Figure 6-64.

Click the “Start” button to start inspiratory valve calibration, the remaining procedure is the same as the pressure sensor calibration.
(6) **Expiratory Valve Calibration**: Click “Expiratory Valve” to enter the interface. A message displayed: “This step is to calibrate the expiratory valve. Please connect patient circuit and test lung before calibration”, as shown in Figure 6-65.

Click the “Start” button to start expiratory valve calibration, the remaining procedure is the same as the pressure sensor calibration.

(7) **Touch Screen Calibration**: Click “Touch screen” to enter the calibration interface. A message is displayed: “This step is to calibrate the touch screen. The ventilator’s screen will disappear during the calibration. Please follow the instruction in the calibration program.” as shown in Figure 6-66.

⚠️ **CAUTION**: Please calibrate the touch screen periodically or when it works abnormally.
Click the "Start" key to start, the remaining procedure is as described in steps above.

(8) Leakage Test Calibration: Click "Leakage Test" to enter the test interface. A message is displayed: "This step is to test the internal leakage of ventilator. Please connect the Insp. Port and Exp. Port directly with a tube", as shown in Figure 6-67.

Click "Start" to start the leakage test. A progress bar and a message "Test in progress, please wait" will be displayed as shown in Figure 6-68. After test, the result will appear: Test succeeded or Test failed. If failed, restart the test.
(9) **Breath Circuit Test Calibration:** Click "Breath Circuit Test" to enter the test interface. A message is displayed: “This step is to test the compliance and leakage of breathing circuit. Please connect the patient circuit to the T-piece, and plug up the patient end of the T-piece”, as shown in Figure 6-69.

Before starting the test, ensure the patient circuit has been connected to the T-piece and the patient end of the T-piece has been plugged up. Click the “Start” button to start the breath circuit test, the remaining procedure is the same as the leakage test.
6.3.6.3.2 Event/Alarm Log

See Section 6.3.5 for detailed description.

6.3.6.3.3 Machine Information

Click the "Machine information" key to enter the Machine information page, this area includes the following information, as shown in Figure 6-70.

1. Software Version:
   a. UI
   b. BDU
   c. Power Supply
2. Runtime Hours
3. O₂ Source Pressure
4. Atmospheric Pressure

![Main Menu](image)

Figure 6-70

6.3.6.3.4 Language

Click the Language button to enter the language screen. English and other languages are available for the user to choose.

6.4 Operation of Other Shortcut Keys

The ventilator has 11 shortcut keys: Inspiratory Hold, Expiratory Hold, Nebulizer, Manual, Suction, Print Screen, Freeze, Screen Lock, Alarm limits, Main Menu and Start
6.4.1 Inspiratory Hold

Inspiratory hold is available within the period of mandatory ventilation and in all modes except full spontaneous breath modes as SPONT/CPAP+PSV, NIV/CPAP.

Press the Inspiratory Hold key during the Inspiratory phase. The operation becomes active when a message stating “Inspiratory Hold” appears with a countdown timer. Keep pressing the Inspiratory Hold key. The expiratory phase will not start until the key is released or after 30 seconds, whichever comes first. If the button is not released after 30 seconds, the system will go to Expiratory state automatically and display a message “Inspiratory hold interrupted!”, as shown in Figure 6-72.

![Image of Inspiratory Hold function on a medical device screen]

Figure 6-71
6.4.2 Expiratory Hold

Expiratory hold is available in all modes. In Expiratory phase, press the Expiratory hold button and the expiratory operation will become active when a message “Expiratory Hold” appears with a countdown timer. The ventilator will stay in the expiratory phase and not transition to the inspiratory phase until either 1) the key is released or 2) 30 seconds have elapsed.

When selected during the expiratory phase, the ventilator will stay in the expiratory phase and not transfer to the inspiratory phase until either 1) the key is released and the current expiratory phase is completed or 2) 30 seconds have elapsed.

Pressing the Expiratory Hold key for more than 30 seconds will cause a message to be displayed. See example in Figure 6-73. Only one expiratory hold will be produced per key press. When Inspiratory hold key is released or if the key is not released after 30 seconds, the key background will revert to normal.

6.4.3 Nebulizer

The nebulizer function is available in all ventilation modes. Press the Nebulizer key turning the key background color to yellow. Meanwhile, a low level alarm “Nebulizer On” is displayed and the message “Nebulizer On, MM min SS s” with countdown timer is displayed in the message area, as shown in Figure 6-74.

Nebulizer flow may be provided by high-pressure O2, and the flow rate is 6L/min±1L/min. To ensure the delivery of tidal volume, the nebulizer is switched off when inspiratory flow rate is less than 15L/min.

The nebulizer will start as inspiratory starts, and the nebulizer will last for the whole inspiratory cycle. When Ventilation mode changes or if the flow is less than 15 L/min, the nebulizer
operation will be interrupted and a medium level alarm “Nebulizer Interrupted” will replace “Nebulizer On” alarm and the countdown timer will stop.

⚠️ **CAUTION**: During nebulization, please connect the filter in front of the expiration valve to prevent the nebulization drug from damaging the expiration flow sensor; inspect, clean and replace the filter regularly.

The nebulizer operation will be cancelled by touching the nebulizer key for longer than 3 seconds in all ventilation modes. The user has two options to clear the “Nebulizer Interrupted” alarm:

1) Remove the source of the shutdown and press the Nebulizer key to restart the Nebulizer operation. Low level alarm “Nebulizer On” will replace “Nebulizer Interrupted” alarm and the message countdown timer will continue or;

2) Acknowledge the alarm by pressing the Alarm Silence key. Nebulizer operation is cancelled and the “Nebulizer Interrupt” alarm is cleared. Pressing the Nebulizer key during a “Nebulizer Interrupted” condition without removing the source of the shutdown will result in the Nebulizer not restarting.

⚠️ **CAUTION**: The ventilator accuracy can be affected by the gas added by use of a nebulizer.

⚠️ **NOTE**: Nebulizer function is not suggested when breath rate is less than 12, to ensure tidal volume delivery.

⚠️ **NOTE**: To ensure the flow volume accuracy, the nebulizer function is disabled if the flow rate of the delivered breath is less than 15 LPM.

⚠️ **CAUTION**: Nebulizer use is not available in noninvasive (NIV) modes.

### 6.4.4 Manual

Manual Trigger is available in all ventilation modes. Press the manual trigger key to initiate a manual breath control as follows, as shown in Figure 6-75.

- For VCV, PRVC, SIMV (VCV): set Vt and Tinsp, to control ventilation.
- For BIVENT: set P\_high and P\_low, to control ventilation switchover.
- For PCV, SIMV (PCV): Set P\_insp, to control ventilation.
- For modes or breath phase with PSV, set P\_supp, to control ventilation.
Manual trigger is also available during backup ventilation period. When initiated during backup mode, the ventilator will remain in backup mode.

The process of Suction support is performed in 3 phases as follows:
• Before aspiration of sputum – 3 minutes of increased oxygen concentration in preparation of airway disconnection;

• Suction Phase - Airway disconnection for suction.

• Post aspiration of sputum – 2 minutes of increased oxygen concentration after reconnection of the airway.

6.4.6 Print Screen

Print Screen shortcut soft key is between the Suction and the Freeze shortcut soft keys, as shown in Figure 6-78. Print Screen is available after machine boot up is finished.

The Print Screen function is intended to be used with a USB Memory device attached to the back of the monitor. With a memory device attached when the Print Screen shortcut softkey is pressed, the system saves a copy of the screen into a JPG file and stores it on the USB device. Each subsequent press of the Print Screen shortcut softkey will create a new JPG file with a unique filename. If the memory stick is full, a message will be displayed stating this.

If the Print Screen shortcut soft key is pressed without a USB memory device connected, the JPG file is saved until a USB memory device is inserted. One and only one screen capture is saved when there is no memory device attached to the monitor.
6.4.7 Freeze

Press “Freeze” key and the current real-time waveforms and loops freeze simultaneously in the main screen when waveform drawing is completed. See example in Figure 6-79. Turing the encoder knob moves the cursor over each point of the waveform and the corresponding measured value is displayed. Pressing Freeze again will restart the waveform and any information displayed during waveform freeze will disappear, and the Freeze key color returns to normal. Waveform Freeze will automatically end 3 minutes after touching the freeze key.

![Figure 6-79](image)

6.4.8 Screen Lock

Press the “Screen Lock” button, displaying a message in the message prompt area. Press and hold Screen Lock for 3 seconds to lock the screen, as shown in Figure 6-80. 3 seconds later, the background color of Screen lock key will turn to yellow and all keys on the touch screen are locked. Instead of the pre-message, a new message is displayed in the message prompt area “Screen Locked. Press Screen Lock for 3 seconds to unlock the screen as shown in Figure 6-81.
Figure 6-80

Figure 6-81
Holding for 3 seconds, the screen lock is cleared, the background color of Screen Lock key turns to normal and a message "Screen Unlocked" is displayed in message prompt area as shown in Figure 6-82.

6.4.9 Alarm Limits

Press the “Alarm limits” button to enter the alarm limits page in the main menu, as shown in Figure 6-83. For detail information and operation refer to Section 6.3.2.
6.5 Ventilation Parameter Set-up

**NOTE:** If ventilation parameter setup keys are not visible along bottom of screen, close the main menu by pressing Main Menu key. See ventilation parameter setup in the main screen as shown in Figure 6-84.

Press the parameter key, turning it yellow. Rotate the encoder knob left or right to modify the parameter value. Press the parameter key again or press the encoder knob to confirm the modified value. If any other area on the screen is touched before confirmation, the new parameter selection is canceled and the previous value will be displayed. As shown in Figure 6-85.
When ventilation mode is changed, the values of the parameter keys displayed will change to correspond to the new ventilation mode.

⚠️ CAUTION: If the encoder knob is not pressed for confirmation, the previous value will be displayed.

The following conditions should be noticed in parameter setting:

a) In order to make parameter setting more safe and reasonable, interlock mode is used for tidal volume $V_T$, ventilator rate and inspiratory time $T_I$ setting. In case any one of the three parameters cannot be adjusted to required value within limits, modify the other two first.

b) Pressure parameter setting is subject to high pressure limit.

c) $P_{SUPP}$ and $P_{INS}$ are relative pressure to PEEP.
6.6 Turn off the Ventilator

(1) Disconnect the breathing hoses from the patient.

(2) Return back to Standby by pressing the Standby key for more than 3 seconds.

(3) Turn off the power switch

(4) Disconnect the gas supply.

(5) Disconnect the power cord from the power supply

⚠️ **NOTE:** Detachable power cord is a means to isolate circuits electrically from AC supply on all poles simultaneously.
7 Alarms and Troubleshooting

WARNING: Only authorized person is permitted to perform maintenance.

7.1 Alarms

CAUTION: In case of alarm, monitor and support the patient first, then carry out troubleshooting later.

An alarm message will appear in alarm information area and alarm indicator will light up. The different colors of alarm in alarm area indicate different priority levels: red is high level (!!!, red indicator flash), yellow is medium (!!, yellow indicator flash) or low level (!, yellow indicator constantly lit).

The operator may be positioned anywhere around the unit to view the alarm light. The alarm light is visible from a distance of 3 meters. To observe the alarm messages the operator position must be in front of the display and within a distance of 1 meter.

7.2 Alarm Message Table

CAUTION: Monitor and support the patient first in case of alarm and perform troubleshooting later.

CAUTION: Operation instructions are not included in the table.

CAUTION: Except for the normal alarm settings, other default alarm settings are changed only by changing the control program and restricted access to changing or to the storage of changes.

CAUTION: If alarm occurs, protect patient safety first, and then go to diagnose fault or service it necessarily.

WARNING: Never leave patient unattended when alarm silence is activated.
**NOTE:** In the table below, L means low level, M means medium level, and H means high level; for the same level, the bigger the alarm number, the higher the alarm level, such as L2 level is higher than L1 level.

<table>
<thead>
<tr>
<th>Alarm Message</th>
<th>Priority</th>
<th>Type</th>
<th>Cause</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>High Airway Pressure</td>
<td>L1</td>
<td>Physiological</td>
<td>Airway Pressure exceeds the setting limit within one ventilation. When the airway pressure exceeds the setting value, the inspiratory phase changes into the expiratory phase within 200 ms and the pressure shall be reduced below PEEP level.</td>
<td>Airway pressure is lower than the alarm limit for three consecutive cycles or at most 15sec.</td>
</tr>
<tr>
<td>Leakage</td>
<td>L2</td>
<td>Physiological</td>
<td>The measured minute volume leak $MV_{LEAK}$ is higher than the minute volume measured on the expiration path for 3 consecutive ventilation cycles or for 10 seconds.</td>
<td>The measured minute volume leak $MV_{LEAK}$ is in normal range for 3 consecutive breaths or for 10 seconds.</td>
</tr>
<tr>
<td>Low Oxygen Supply</td>
<td>L3</td>
<td>Physiological</td>
<td>Oxygen source is lower than 160kPa $O_2$ is set to 21%.</td>
<td>Oxygen supply is above 160kPa.</td>
</tr>
<tr>
<td>AC Failure</td>
<td>L4</td>
<td>Technical</td>
<td>During ventilator operation, when an AC power failure occurs and there is no battery power, the power board will alarm for 120 seconds minimum. When powered from batteries, an “AC Failure” alarm would occur.</td>
<td>During ventilator operation, when an AC power failure occurs and there is no battery power, the power board will alarm for 120 seconds minimum. When powered from batteries, an “AC Failure” alarm would occur. When battery is supplied, produces a low-level alarm and gives the prompt message of “AC power faulty” alarm.</td>
</tr>
<tr>
<td>Alarm Message</td>
<td>Priority</td>
<td>Type</td>
<td>Cause</td>
<td>Remedy</td>
</tr>
<tr>
<td>----------------------------------</td>
<td>----------</td>
<td>---------------</td>
<td>----------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------</td>
</tr>
<tr>
<td>Nebulizer On</td>
<td>L5</td>
<td>Physiological</td>
<td>Start Nebulizer operation</td>
<td>Nebulizer operation is completed or Interrupted.</td>
</tr>
<tr>
<td>Int. Battery Calibration required</td>
<td>L6.1</td>
<td>Technical</td>
<td>During ventilator operation, when an AC power failure occurs and there is the int. battery power, an “Int. Battery Calibration required” alarm occurs. When the ventilator starts running and there is the int. battery power, an “Int. Battery Calibration required” alarm occurs.</td>
<td>Calibrate the int. battery. Then restart the ventilator.</td>
</tr>
<tr>
<td>Opt. Battery Calibration required</td>
<td>L6.2</td>
<td>Technical</td>
<td>During ventilator operation, when an AC power failure occurs and there is the opt. battery power, an “Opt. Battery Calibration required” alarm occurs. When the ventilator starts running and there is the opt. battery power, an “Opt. Battery Calibration required” alarm occurs.</td>
<td>Calibrate the opt. battery. Then restart the ventilator.</td>
</tr>
<tr>
<td>Expiratory Hold Interrupted</td>
<td>L7</td>
<td>Physiological</td>
<td>Start Expiratory holding</td>
<td>Expiratory hold operation is completed.</td>
</tr>
<tr>
<td>Inspiratory Hold Interrupted</td>
<td>L8</td>
<td>Physiological</td>
<td>Start the inspiratory holding</td>
<td>Inspiratory holding operation is completed.</td>
</tr>
<tr>
<td>Low Battery</td>
<td>M1</td>
<td>Technical</td>
<td>Under the battery operation, the remaining battery run time is less than 30min.</td>
<td>When “Limited Battery Capacity” alarm appears or AC power supply starts.</td>
</tr>
<tr>
<td>Alarm Message</td>
<td>Priority</td>
<td>Type</td>
<td>Cause</td>
<td>Remedy</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>----------</td>
<td>------------</td>
<td>----------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------</td>
</tr>
<tr>
<td>High Respiration Rate</td>
<td>M2</td>
<td>Physiological</td>
<td>Respiration Rate has exceeded the set limit for four consecutive ventilation cycles or continuously for 20sec.</td>
<td>Respiration Rate is less than the set limit for four consecutive ventilation cycles.</td>
</tr>
<tr>
<td>Oxygen Sensor Failure</td>
<td>M3</td>
<td>Physiological</td>
<td>Checked during pre-inspection before use.</td>
<td>Measure signals of the sensor through equipment inspection.</td>
</tr>
<tr>
<td>High Oxygen Concentration</td>
<td>M4</td>
<td>Physiological</td>
<td>Oxygen concentration exceeds the preset oxygen concentration +6% continuously for 30sec.</td>
<td>Oxygen concentration is lower than the preset oxygen concentration + 6% for 30sec continuously.</td>
</tr>
<tr>
<td>Low Oxygen Concentration</td>
<td>M5</td>
<td>Physiological</td>
<td>Oxygen concentration is lower than the preset oxygen concentration -6% or lower than 18% continuously for 30sec.</td>
<td>Oxygen concentration is higher than the preset oxygen concentration - 6% or higher than 18% for 30sec continuously.</td>
</tr>
<tr>
<td>High Expiratory Minute Volume</td>
<td>M6</td>
<td>Physiological</td>
<td>Minute Volume exceeds the set limit, after 1min of ventilation</td>
<td>Minute Volume lower than the set limit for 15sec.</td>
</tr>
<tr>
<td>Low Expiratory Minute Volume</td>
<td>M7</td>
<td>Physiological</td>
<td>Minute Volume is lower than the set limit within 1min.</td>
<td>Minute Volume higher than the set limit for 15sec.</td>
</tr>
<tr>
<td>Alarm Message</td>
<td>Priority</td>
<td>Type</td>
<td>Cause</td>
<td>Remedy</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>----------</td>
<td>------------</td>
<td>----------------------------------------------------------------------</td>
<td>------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Maximum Inspiratory Time</td>
<td>M8</td>
<td>Physiological</td>
<td>NIV – inspiration duration exceeds Tispont setting for three times in succession</td>
<td>Inspiratory period is normal under NIV mode.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Invasive - inspiration duration exceeds 5 seconds for adult and 2 seconds for child for three times in succession</td>
<td>Within the specified condition for two consecutive PSV ventilation under invasive mode.</td>
</tr>
<tr>
<td>Nebulizer Interrupted</td>
<td>M13</td>
<td>Physiological</td>
<td>Nebulizer operation has been interrupted by mode changes.</td>
<td>Nebulizer operation continued or alarm is acknowledged by pressing Alarm Silence key.</td>
</tr>
<tr>
<td>Low Inspiratory Tidal Volume</td>
<td>M14</td>
<td>Physiological</td>
<td>Inspiratory Tidal Volume exceeds the upper limit for one ventilation.</td>
<td>Inspiratory Tidal Volume is lower than the alarm limit for one ventilation.</td>
</tr>
<tr>
<td>Limited Battery Capacity</td>
<td>H1</td>
<td>Technical</td>
<td>Under the battery operation, the remaining battery run time is less than 10min.</td>
<td>AC power supply starts.</td>
</tr>
<tr>
<td>Internal Error</td>
<td>H2.1</td>
<td>Technical</td>
<td>Internal computing error</td>
<td>Cycle AC Power-if alarm occurs again stop using Ventilator, note code displayed and call for Service.</td>
</tr>
<tr>
<td>Communication Error BDU</td>
<td>H2.2</td>
<td>Technical</td>
<td>Failure detected in BDU communications.</td>
<td>Cycle power. If the alarm occurs multiple times, call for Service.</td>
</tr>
<tr>
<td>Alarm Message</td>
<td>Priority</td>
<td>Type</td>
<td>Cause</td>
<td>Remedy</td>
</tr>
<tr>
<td>----------------------------</td>
<td>----------</td>
<td>------------</td>
<td>----------------------------------------------------------------------</td>
<td>------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Communication Error PS</td>
<td>H2.3</td>
<td>Technical</td>
<td>Failure detected in PS communications.</td>
<td>Cycle power. If the alarm occurs multiple times, call for Service.</td>
</tr>
<tr>
<td>S/W Mismatch Error</td>
<td>H2.4</td>
<td>Technical</td>
<td>System detects software versions are incorrect.</td>
<td>Call Service.</td>
</tr>
<tr>
<td>Fan Blocked</td>
<td>H2.6</td>
<td>Technical</td>
<td>System detects cooling fan failure</td>
<td>Remove any obstructions. Cycle Power. If the alarm occurs again, call Service.</td>
</tr>
<tr>
<td>BDU Failure</td>
<td>H2.7</td>
<td>Technical</td>
<td>Internal hardware failure detected.</td>
<td>Check to make sure inspiratory and expiratory circuits have no obstruction. Retry power-up tests. If this alarm persists, Call Service.</td>
</tr>
<tr>
<td>Continuous Airway Pressure</td>
<td>H3</td>
<td>Physiological</td>
<td>Airway pressure has exceeded PEEP+15cmH$_2$O continuously for 15sec. Expiratory valve is opened for gas release.</td>
<td>Airway pressure is lower than PEEP+15cmH$_2$O continuously for 5sec.</td>
</tr>
<tr>
<td>High Airway Pressure</td>
<td>H4</td>
<td>Physiological</td>
<td>Airway Pressure is lower than the set limit within three consecutive ventilation cycles.</td>
<td>Airway pressure exceeds the alarm limit for three consecutive cycles or at most 15sec.</td>
</tr>
<tr>
<td>Alarm Message</td>
<td>Priority</td>
<td>Type</td>
<td>Cause</td>
<td>Remedy</td>
</tr>
<tr>
<td>-----------------------</td>
<td>----------</td>
<td>---------------------</td>
<td>----------------------------------------------------------------------</td>
<td>------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Apnea</td>
<td>H5</td>
<td>Physiological</td>
<td>No ventilation cycles within the apnea period.</td>
<td>Refer to Chapter 11.1.3 for Apnea resetting details.</td>
</tr>
<tr>
<td>Low Expiratory Tidal Volume</td>
<td>H6</td>
<td>Physiological</td>
<td>Expiratory Tidal Volume is lower than the setting value for three consecutive ventilation cycles or continuously for 30s.</td>
<td>Expiratory Tidal Volume is higher than the setting value for three consecutive ventilation cycles or continuously for 30s.</td>
</tr>
<tr>
<td>High PEEP</td>
<td>H7</td>
<td>Physiological</td>
<td>PEEP measurement is higher than the alarm limit for three consecutive ventilation cycles.</td>
<td>PEEP is lower than the alarm limit for three consecutive cycles or at most 15sec.</td>
</tr>
<tr>
<td>Circuit Disconnect</td>
<td>H8</td>
<td>Physiological</td>
<td>Ventilating circuit is disconnected.</td>
<td>Re-connect patient circuit.</td>
</tr>
<tr>
<td>Occlusion</td>
<td>H9</td>
<td>Physiological</td>
<td>Ventilating circuit is occluded.</td>
<td>Occlusion is removed.</td>
</tr>
<tr>
<td>Leakage out of range</td>
<td>H10</td>
<td>Physiological</td>
<td>NIV Mode only- Leak exceeds 75% of the maximum volume compensation capacity.</td>
<td>Leak is within 75% of the maximum volume compensation capacity.</td>
</tr>
<tr>
<td>Low Airway Pressure</td>
<td>H11</td>
<td>Physiological</td>
<td>Airway pressure is lower than the set limit for three consecutive ventilation cycles.</td>
<td>Airway pressure is higher than the alarm limit for three consecutive ventilation cycles or at most 15 sec.</td>
</tr>
<tr>
<td>Low Oxygen Supply</td>
<td>H12</td>
<td>Physiological</td>
<td>Oxygen supply is lower than 160kPa.</td>
<td>Oxygen supply is above 160kPa.</td>
</tr>
<tr>
<td>Alarm Message</td>
<td>Priority</td>
<td>Type</td>
<td>Cause</td>
<td>Remedy</td>
</tr>
<tr>
<td>-------------------</td>
<td>----------</td>
<td>------------</td>
<td>----------------------------------------------------------------------</td>
<td>------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Low PEEP</td>
<td>H13</td>
<td>Physiological</td>
<td>PEEP measurement is lower than the alarm limit for three consecutive ventilation cycles.</td>
<td>PEEP is higher than the alarm limit for three consecutive cycles or at most 15sec.</td>
</tr>
<tr>
<td>CO₂ Accuracy Error</td>
<td>H14</td>
<td>Physiological</td>
<td>The CO₂ module is not detected during the probation period or the CO₂ module fails.</td>
<td>Redetect or replace the CO₂ module.</td>
</tr>
<tr>
<td>CO₂ Adapter Failure</td>
<td>H15</td>
<td>Physiological</td>
<td>Detected by CO₂ Sensor.</td>
<td>Reconnect the CO₂ Adapter. If alarm occurs again, replace the CO₂ Adapter.</td>
</tr>
<tr>
<td>CO₂ sensor Comm. Failure</td>
<td>H16</td>
<td>Physiological</td>
<td>The communication between the CO₂ module and the host instrument fails.</td>
<td>Check the cable between the CO₂ module and the host instrument.</td>
</tr>
<tr>
<td>CO₂ Sensor Error</td>
<td>H17</td>
<td>Physiological</td>
<td>Detected by CO₂ Sensor.</td>
<td>Reconnect the CO₂ sensor. If alarm occurs again, replace the CO₂ sensor.</td>
</tr>
<tr>
<td>Low EtCO₂</td>
<td>H18</td>
<td>Physiological</td>
<td>EtCO₂ is lower than the setting value.</td>
<td>Increase EtCO₂ or reduce the setting value.</td>
</tr>
<tr>
<td>High EtCO₂</td>
<td>H19</td>
<td>Physiological</td>
<td>EtCO₂ is higher than the setting value.</td>
<td>Reduce EtCO₂ or increase the setting value.</td>
</tr>
</tbody>
</table>
8 User Maintenance

⚠️ **WARNING:** Only authorized person is permitted to perform maintenance.

⚠️ **WARNING:** Our company recognizes that cleaning, disinfection, and sterilization practices vary widely among medical institutions. It is not possible to specify or require specific practices that will meet all needs, or to be responsible for the effectiveness of cleaning, sterilization, and other practices carried out in the patient care setting.

⚠️ **WARNING:** Use a cleaning and disinfection schedule that conforms to your institution’s disinfection and risk-management policies:

- Refer to the material safety data as applicable.
- Refer to the operation and maintenance manuals of all disinfection equipment.
- Do not inhale fumes that may result from any disinfection process.

⚠️ **WARNING:** Movable and removable parts may clamp or even crush your hand. Use caution when moving or replacing system components.

⚠️ **CAUTION:** The disposal of environmentally harmful devices (such as batteries and LCD display) must be in accordance with local regulations.

⚠️ **WARNING:** Do not use talc, zinc stearate, calcium carbonate, corn starch or similar material to prevent sticking of the bellows, as these materials may enter the patient’s lungs or airway, causing irritation or injury.

⚠️ **CAUTION:** To prevent system damage:

- Refer to the literature supplied by the manufacturer of the cleaning agent.
- Never use organic, halogenated or petroleum-based solvents, anesthetic, glass cleaning agents, acetone or other irritant agents.
- Never use abrasive agents (i.e. steel wool or silver polish) to clean components.
- Keep all liquids away from electronic components.
- Prevent liquid from entering the equipment.
- All cleaning solutions used must have a pH between 7.0 and 10.5.
CAUTION: Never immerse the oxygen sensor or its connector in any type of liquid.

- Dispose of the oxygen sensor per the manufacturer’s specification.

CAUTION: Do not wash the inner surface of the oxygen sensor.

CAUTION: Prior to use after cleaning or disinfecting, power up the system as described in section 6 and follow the on-screen Pre-Use test prompts to perform the Leak Test and Circuit Compliance Test.

8.1 Cleaning and Disinfection

CAUTION: Before the first use clean, disinfect, and sterilize the ventilator. Disposable components must be disposed in accordance with local regulations. Don’t use hard brushes or other sharp tools in cleaning to avoid damage to parts.

8.1.1 Cleaning and Disinfecting Agents/ Autoclaving

<table>
<thead>
<tr>
<th>Agent</th>
<th>Classification</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Mild dishwashing detergent</td>
<td>Detergent</td>
</tr>
<tr>
<td>• Soapy water with detergent ph between 7.0 and 10.5</td>
<td>Detergent</td>
</tr>
<tr>
<td>• Isopropyl alcohol (70% solution)</td>
<td>Intermediate level disinfectant</td>
</tr>
<tr>
<td>• Window cleaning solution (with isopropyl alcohol and ammonia)</td>
<td>Intermediate level disinfectant</td>
</tr>
<tr>
<td>• Sodium hypochloride- (bleach) in water (10% solution)</td>
<td>Intermediate level disinfectant</td>
</tr>
<tr>
<td>• Hydrogen peroxide (3% solution)</td>
<td>Intermediate level disinfectant</td>
</tr>
<tr>
<td>• Gluteraldehyde 2% solution</td>
<td>High level disinfectant</td>
</tr>
<tr>
<td>• Steam autoclaving up to a maximum temperature of 134°C (273°F).</td>
<td>High level disinfectant</td>
</tr>
</tbody>
</table>
8.1.2 Cleaning and Disinfection Methods

Different parts of the ventilator have their respective cleaning and disinfection methods. The following categories are defined for the parts noted in Table 8-1. The parts need to be cleaned, disinfected and thoroughly dried before reassembly.

A: Wipe: If there is a potentially infectious substance on the breathing system, such as blood or secretion, wipe away the substance with disposable cloth using proper disinfectant. Use a soft cloth with a water-soluble detergent or disinfectant wipes.

B: Machine washing: Automatic washing with washer and disinfecting with disinfection machine.

C: Immersion disinfection: Soak in glutaraldehyde-based formulations of 2%.

D: High temperature and pressure disinfection: At 121°C for 20 minutes minimum, or at 134°C for 8 minutes minimum. Follow the manufacturer’s instructions for high level disinfection.

High-temperature disinfection does not have any cleaning effect. It should only be used on components that have already been cleaned by hand or machine and then thoroughly dried.

Table 8-1

<table>
<thead>
<tr>
<th>Part name</th>
<th>Cleaning and disinfection methods</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ventilator exterior, including housing, gas supply hoses and power cord</td>
<td>✓</td>
</tr>
<tr>
<td>All components in Figure 8-3</td>
<td>✓ ✓ ✓ ✓</td>
</tr>
<tr>
<td>CO₂ sensor, other breathing circuit parts or accessories</td>
<td>Follow the manufacturer’s guidelines</td>
</tr>
</tbody>
</table>

8.1.3 Cleaning and Disinfection of Components

8.1.3.1 External Surfaces

Using a soft cloth with a water-soluble detergent or disinfectant wipes, clean the housing, gas supply hoses and power cord.

8.1.3.2 Expiratory Module

(1) Disassembly
To remove the components from expiratory module:

a. Press the button (1 in Figure 8-1);

b. Pull out the expiratory valve core component (2 in Figure 8-1, Figure 8-2);

c. Remove the end cover (1 in Figure 8-3) and the gasket (2 in Figure 8-3);

d. Remove the scale board (3 in Figure 8-3) and the diaphragm (4 in Figure 8-3);

e. Rotate the expiratory connector (5 in Figure 8-3) clockwise;

f. Pull out one-way valve core (6 in Figure 8-3), rubber o-sealing ring (7 in Figure 8-3) and one-way diaphragm (8 in Figure 8-3);

g. Rotate the water trap (9 in Figure 8-3), then pull it out.

(2) Cleaning

a. Wash each component using a mild detergent and water solution.
b. Rinse with clean, hot water and allow to thoroughly dry.

(3) Disinfection

⚠️ **NOTE:** Ensure that all the components have been cleaned before disinfecting.

Using the Gluteraldehyde disinfection solution, follow the manufacturer’s instructions for high level disinfection and rinsing of all components while adhering to facility policies and procedures.

All the components can also be high temperature and pressure disinfected. Using an autoclave, follow the manufacturer’s instruction for high level disinfection of all the components while adhering to facility procedures.

(4) Assembly

Reassemble the components in the reverse order.

After connection, please perform a pre-use test and verify all tests passed.

---

### 8.2 Regular Maintenance

#### 8.2.1 Maintenance Principles

Do not use a faulty machine. Ask an authorized agent of our company to carry out all necessary maintenance tasks. Test the machine after maintenance for normal operation. Every parameter should meet requirement in specification.

In order to ensure the reliability of the machine, all maintenance and repair work should be carried out by an authorized agent of our company.

⚠️ **CAUTION:** Only authorized personnel are permitted to perform maintenance.

Use products of our company to replace damaged ones and test, ensuring all specifications met.

Contact local service agent of our company in case support is needed. In all cases, maintenance fee is the current component price plus reasonable labor cost, except for those within guarantee period.

#### 8.2.2 Periodic Maintenance Schedule

The schedule is designed based on the typical condition, that is to say, the least maintenance times vs 2000h operating per year. In case the actual operating time is longer than 2000h per year, the maintenance times should be more.
If necessary, our company can provide circuit diagram, service part list, calibration instructions to assist authorized service personnel.

<table>
<thead>
<tr>
<th>Minimum maintenance interval</th>
<th>Task</th>
</tr>
</thead>
<tbody>
<tr>
<td>Daily</td>
<td>Drain water in gas supply inlet filter, check liquid in expiratory module water trap (collected liquid volume cannot be more than half of the bottle).</td>
</tr>
<tr>
<td>Weekly</td>
<td>Calibrate oxygen sensor</td>
</tr>
<tr>
<td>1-3 month(s)</td>
<td>Clean air filter Clean cooling fan filter on rear of machine.</td>
</tr>
<tr>
<td>Every year</td>
<td>Calibrate flow sensor and pressure sensors; Calibrate inspiratory valve and expiratory valve (if necessary).</td>
</tr>
<tr>
<td>Every 6 months</td>
<td>Charge and discharge the batteries once (Charge time: at least 3.5h).</td>
</tr>
<tr>
<td>Every year or after a failed calibration</td>
<td>Replace the O₂ sensor (actual life depends on temperature and O₂ concentration)</td>
</tr>
<tr>
<td>After cleaning and connecting</td>
<td>Check components and replace or repair when necessary.</td>
</tr>
</tbody>
</table>

⚠️ **WARNING:** If the Ventilator will not be in use for a period of more than 6 months, the internal batteries must be disconnected or removed to prevent possible damage to the equipment or risk to users or service personnel.

⚠️ **WARNING:** The ventilator must not use, nor be connected to, any anti-static or electrically conductive hoses and tubing.

### 8.2.3 Service Life of Product/Accessories

Service Life is defined as the time that the manufactured device can be expected to be ‘serviceable’ or supported by our company and the maximum time the device can be used safely. Some items within the device may require maintenance, repair or replacement within this time. Such items will be available from our company for the service life of the product. The calculation of the service life begins at the connection of the product at the customer site. Our company recommends for safe use that each device is replaced after its service life is completed.

⚠️ **CAUTION:** The service life of the following items is based on specified operating conditions.
### 8.3 Maintenance in Operation and Transportation

The location of machine should be proper so that it cannot obstruct or be disturbed by medical care personnel. Fix power supply cable well to avoid failure. Use caution not to touch accidental keys on the panel, which may make tidal volume setting wrong.

During transportation of the ventilator, with or without a patient connected, make sure that the following conditions are fulfilled:

- Gas cylinders are connected with a sufficient amount of gas and the Battery module is functioning. Follow the hospital guidelines.
- Use the handles on the Mobile Cart. Transport the bed and the ventilator slowly, and watch the patient connection carefully to see that no pulling or other movement occurs.
- Be careful not to tip the Mobile Cart when crossing an obstacle such as a doorstep.

#### 8.3.1 Transportation

Use care when moving machine within hospital or clinical environment.

⚠️ **WARNING:** If Control Unit of Ventilator is dropped or damaged during transportation, equipment failure could result in patient injury.

⚠️ **WARNING:** Tip over hazard – use care when moving Ventilator mounted to Cart as device could tip over leading to injury or damage to the equipment and possible subsequent patient injury.

User can carry packaged machine while riding in vehicle, plane and train. Impact, severe shock and moisture should be avoided during transportation, with ambient temperature -20°C~+60°C and relative humidity not more than 95%. In case transportation conditions do not meet this requirement, put the machine in specified operating environment for at least 24h before using.
8.3.2 Storage

⚠️ **CAUTION:** Do not put ventilator into the shock environment.

⚠️ **CAUTION:** Do not lay heavy items on the top of the ventilator.

The machine should be stored in a room with temperature -20°C to + 60°C and relative humidity not more than 95% non-condensing, with ventilation and no corrosive gases.

⚠️ **WARNING:** If the storage environment conditions don't agree, put the machine in specified operating environment at least 24h before using.

⚠️ **CAUTION:** The device should be stored at the room that is ventilated and in which no corrosive gases exist.

⚠️ **CAUTION:** When the storage conditions are beyond the requirements of operational environment, and the storage state is transferred into operation state, the product only can be used after being stored in environment for over 24 hours.

8.4 Consumables Replacement

8.4.1 Fuse Replacement

⚠️ **WARNING:** Before replacing fuse, first disconnect AC power. Otherwise, it will cause injury or even death.

⚠️ **WARNING:** When replacing fuse, make sure the new fuse is the same type and size as the old one; otherwise, the ventilator will be damaged.

⚠️ **CAUTION:** Fuse is a damageable part, which shall be replaced with moderate force and speed.

1. **AC Circuit Fuse**

   **Fuse replacement steps:**
   - Insert screwdriver into the trench (2) of end of fuse box, see Figure 8-4.
   - Pull out fuse holder (1).
   - Remove the fuse (3).
   - Load new fuse.
   - Push the new fuse into the original position gently.
   - Connect AC power, and then start the ventilator to test.
2. **DC Power Fuse**

   **Fuse replacement steps:**
   - Insert screwdriver into the trench (1) of end of fuse box, see Figure 8-2.
   - Pull out fuse holder (2).
   - Remove the fuse (3).
   - Load new fuse.
   - Push the new fuse into the original position gently.

---

**8.4.2 Battery Maintenance**

⚠️ **NOTE:** One or more batteries must be used.

⚠️ **NOTE:** Before the machine is put into patient use, the battery must be fully charged. If the battery must not be fully charged and AC power supply fails, be careful about the battery power.
**Battery specification**

Battery module:

-- DC12V, 6.6AH, 14.4V lithium-ion battery  
-- Typical charge time: 3.5 hours  
-- Typical discharge time: 2 hour

When the main power supply voltage is too low or the main power supply fails, two backup batteries (one is necessary, and one is optional) can protect the ventilator. When having a power failure, the ventilator can switch to battery supply automatically, and can normally work without pneumatic power supply failure. The two batteries are usually available for the ventilator working for 4 hours.

**Precautions**

Charge: When operating with AC power supply on, the system will maintain the battery automatically. Charge time is less than 3.5 hours.

Discharge: The machine is operating on battery.

In case of low battery condition, an alarm message “Low battery” will appear, notifying the user to restore AC power supply to charge, otherwise the batteries will be depleted and another alarm “Limited Battery Capacity” will be displayed, and eventually the system will shut down (for safety reason, manual power-on is required to start the machines again after an automatic shutdown).

Before the machine is put into patient use, or disconnect AC power for the transport or other purposes, check the battery power. If the battery is not fully charged, connect the ventilator to AC power for at least 3.5h and recharge the battery until the power reaches 80%~100%.

**Battery storage**

In case the battery is to be stored for a long time, charge it fully prior to storage.

To keep the battery power and prolong the battery life, please ensure that the ventilator is connected to the main power. Charge the battery every six months, while the actual time depends on the storage environment.

High humidity and high temperature environments should be avoided for storage.

**Battery replacement**

Open the battery cover (See Figure 2-5), then remove the battery from its housing.

Same model battery with CE certification is suggested. Make sure AC power supply is disconnected before replacing.
CAUTION: An authorized service representative can replace the battery. If the battery is not to be used for a long-time, please contact the service representative to disconnect the battery. The waste battery should be disposed of in accordance with the local policies.

- **Battery charging and calibration**

  Use the battery charger supplied by our company to charge or calibrate the battery. After calibrating the battery, the ventilator can read the residual battery capacity accurately.

  Please charge or calibrate the battery according to the instructions of the battery charger.

CAUTION: When ‘low battery’ alarm occurs, charging should be done immediately. Otherwise the Ventilator System could shut off automatically in several minutes.

**8.4.3 Oxygen Sensor**

The oxygen sensor can be used to measure the local oxygen concentration when it is connected to the ventilator or other equipments. The oxygen sensor is suitable for adult and child.

**8.4.3.1 Oxygen Sensor Replacement**

**Disassembly**

Step 1: Open outward (1), and then remove the cover.

![Figure 8-6](image)

Step 2: Turn the oxygen sensor (3) anticlockwise, and then remove it.
Assembly

Inspect the oxygen sensor for damage and replace as necessary. Then reassemble the oxygen sensor.

8.4.3.2 Oxygen Sensor Calibration

For oxygen sensor calibration, refer to section 6.8.3.1.

8.4.3.3 Technical Specifications of Oxygen Sensor

<table>
<thead>
<tr>
<th>Specification</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Measure range</td>
<td>0-1500 mBar O₂</td>
</tr>
<tr>
<td>Measure accuracy</td>
<td>&lt;1%</td>
</tr>
<tr>
<td>Response time</td>
<td>&lt;15 s (air to 100% O₂)</td>
</tr>
<tr>
<td>Operating temperature</td>
<td>-20°C ~ +50°C</td>
</tr>
<tr>
<td>Operating pressure</td>
<td>0.5 - 2.0 Bar</td>
</tr>
<tr>
<td>Operating humidity</td>
<td>0 - 99% RH non-condensing</td>
</tr>
<tr>
<td>Storage temperature</td>
<td>0°C to 20°C</td>
</tr>
<tr>
<td>Drift</td>
<td>&lt;200 µV</td>
</tr>
<tr>
<td>Reproducibility</td>
<td>± 1%</td>
</tr>
<tr>
<td>Type</td>
<td>Chemical fuel cell</td>
</tr>
<tr>
<td>Expect life time</td>
<td>0.94 x 10⁶% O₂ hours at 20°C</td>
</tr>
<tr>
<td></td>
<td>0.6 x 10⁶% O₂ hours at 40°C</td>
</tr>
<tr>
<td>Total system response time</td>
<td>&lt;60s</td>
</tr>
<tr>
<td>Working principle of O₂ monitor</td>
<td>The O₂ monitor surveys and displays the O₂ concentration in the patient loop. The oxygen sensor component contains an oxygen sensor, which can produce the voltage proportional to the oxygen partial pressure (concentration) on its detection surface.</td>
</tr>
</tbody>
</table>
The oxygen sensor is an electrochemical device (chemical battery). Oxygen expands in this device through a layer of film and oxidizes the base metal electrode. The oxidation process produces a current with an amplitude proportional to the oxygen partial pressure indicated by the electrode sensor. The base metal electrode is gradually exhausted during the oxidation process.

The voltage of the sensor is influenced by the temperature of the monitoring gas mixture. The surgical thermosensitive resistor of the sensor automatically compensates temperature change in the sensor.

The \( O_2 \) monitor converts the sensor signal into the corresponding oxygen percentage value by using signal processing and circuit analysis. The system displays the value and compares it with the stored alarm limit value. If the value exceeds the threshold, the monitor will alarm.

⚠️ **NOTE**: After being in a condensing atmosphere, the oxygen sensor shall be stored for more than 24 hours in an environment equivalent to operating humidity.

### 8.4.3.4 Oxygen Sensor Maintenance

The oxygen sensor should be regularly calibrated. For the calibration interval, refer to section 8.2.2.

To improve the life time of the oxygen sensor, when the ventilator is not in use the oxygen sensor should be avoided contact with the high-concentration oxygen.

The oxygen sensor is consumptive, and the period of valid is ordinarily 12 months. So the user should pay attention to the valid of the oxygen sensor. When the oxygen sensor fails, please contact the manufacturer.

The recommended oxygen sensor is produced by our company.

⚠️ **WARNING**: Do not immerse oxygen sensor in liquid. Do not conduct autoclave or high temperature fumigation on the oxygen sensor.

### 8.4.4 Paramagnetic Oxygen Sensor (optional)

The paramagnetic oxygen sensor can be used to measure the local oxygen concentration when it is connected to the ventilator or other equipment. The oxygen sensor is suitable for adult and child.
8.4.4.1 Paramagnetic Oxygen Sensor Calibration

For the paramagnetic oxygen sensor calibration, refer to section 6.3.3.1.

8.4.4.2 Technical Specifications of Paramagnetic Oxygen Sensor

<table>
<thead>
<tr>
<th>Performance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Technology</td>
</tr>
<tr>
<td>Range</td>
</tr>
<tr>
<td>Accuracy (Intrinsic Error)</td>
</tr>
<tr>
<td>Linearity</td>
</tr>
<tr>
<td>Repeatability</td>
</tr>
<tr>
<td>Zero Drift</td>
</tr>
<tr>
<td>Response Time (T₁₀ – T₉₀)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Outputs/Controls</th>
</tr>
</thead>
<tbody>
<tr>
<td>Signal Output</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Physical</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weight</td>
</tr>
<tr>
<td>Dimensions</td>
</tr>
<tr>
<td>Diffusion Port</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sample condition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample Gas Condition</td>
</tr>
<tr>
<td>Gas Exchange</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Ambient conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Operation Temperature</td>
</tr>
<tr>
<td>Storage Temperature</td>
</tr>
<tr>
<td>Temperature Coefficient</td>
</tr>
<tr>
<td>Operating Pressure Range</td>
</tr>
<tr>
<td>Ambient Humidity</td>
</tr>
<tr>
<td>External Power Supply</td>
</tr>
<tr>
<td>Power Consumption</td>
</tr>
</tbody>
</table>
8.4.5 Diaphragm Replacement

Diaphragm replacement method:

h. Press the button (1 in Figure 8-8);

i. Pull out the expiratory valve core component (2 in Figure 8-8);

j. Remove the safety valve cover (1 in Figure 8-9) and the gasket (2 in Figure 8-9);

k. Remove the scale board (3 in Figure 8-9) and the diaphragm (4 in Figure 8-9);

l. Reassemble the above components in the reverse order.

Test: When the diaphragm and scale board are replaced, the inspiratory valve must be calibrated.; Also, perform a pre-use test and verify all test pass.
The one-way diaphragm replacement method:

a. Remove the expiratory valve core component;

b. Remove the one-way valve core component (1);

c. Remove one-way diaphragm(2);

d. Reassemble the above components in the reverse order.

Test: When the one-way diaphragm has been replaced, the inspiratory valve must be calibrated.
8.4.7 Fan Filter Cotton Replacement

The replacement method of the fan filter cotton:

a. Remove the fan filter cotton cover(2);

b. Remove the fan filter cotton(1);

c. Reassemble the above components in the reverse order.
8.4.8 Filter Element of Gas Inlet Replacement

Figure 8-12

Filter element of gas inlet replacement method:

a. Remove the filter cover(1);

b. Remove the filter support sleeve(2);

c. Remove the filter element of gas inlet(3);

d. Reassemble the above components in the reverse order.
8.4.9 Filter Replacement (Part No.: 130003930)

The filter replacement method:

a. Remove the two screws (1) (Hexagon socket head cap screws M4×10);

b. Remove the oxygen inlet connector (2);

c. Remove the filter (3);

d. Reassemble the above components in the reverse order.

8.5 Disposal

This product must not be disposed of with your other waste. Instead, it is your responsibility to dispose of your waste equipment by handing it over to a designated collection point for the recycling of waste electrical and electronic equipment, or by returning it to our company for reprocessing. The separate collection and recycling of your waste equipment at the time of disposal will help to conserve natural resources and ensure that it is recycled in a manner that protects human health and the environment. For more information about where you can drop off your waste equipment for recycling, please contact your local city office, your waste disposal service, or your product distributor or retailer.
**WARNING:** Treatment of batteries and O₂ sensor:

- Follow all local regulations with respect to environmental protection when disposing of batteries and O₂ sensor. These products contain toxic compounds irrespective of physical condition. They should be disposed of according to local waste management requirements and environmental legislation. They should not be burned since they may give off toxic fumes.
- Do not throw into fire! Risk of explosion.
- Do not force open! Danger of bodily injury.
9 Warranty

Manufacturing techniques and materials:

For a period of one year from the date of original delivery to the customer, the components and assemblies of this product are warranted to be free from defects caused by manufacturing techniques and materials, provided that the equipment is properly operated under conditions of normal use and that the equipment is regularly maintained per requirements specified by our company. The warranty period for other parts is three months. Consumable parts are not included in this warranty. Our company’s obligation under the above warranty is limited to repairing the equipment free of charge.

Freedom from Obligations:

- Our company’s obligation under the above warranties does not include freight and other fees;
- Our company is not responsible for any direct or indirect product damage or delays which result from improper use; alteration by using assemblies in an unapproved configuration; and maintenance by anyone other than our company or our company’s appointed representatives;
- This warranty does not apply to the following situations:
  - Improper use;
  - Machines used without the proper maintenance or damaged;
  - Our company original serial number label or mark is removed or replaced.

Security, reliability and operating condition:

Our company is not responsible for the security; reliability and operating condition of this product in cases where:

- The assemblies are disassembled, extended and readjusted
- The product is not operated correctly in accordance with these User manual instructions.
- The AC power used or the operating environment does not follow the requirements in this manual.

9.1 Return

Follow these steps in case the product must be returned to our company:

1. Obtain the rights of return (Return Material Authorization Number)
Contact our Customer Service and inform us of the part number and type of the product. The part number is marked on the surface of the product. Return is not allowed if the number cannot be identified. Receive an RMA number from our company and mark it on the exterior of
the product packaging. Enclose a statement of the product number, product type and also the reason for return.

2. Transportation charges
Transportation and insurance charges must be prepaid by the user for transporting the product to our company for repair.
10 Theory of Operation

10.1 Ventilation Modes

There are Invasive and non-invasive (NIV) ventilation modes in the ventilator. All the modes are suitable for children and adults.

⚠️ CAUTION: Nebulizer use is not available in noninvasive (NIV) modes.

Invasive ventilation modes include:

- Assist/Control ventilation (A/C) modes. These modes allow mandatory ventilation and include VCV, PCV and PRVC.
- Synchronous Intermittent Mandatory Ventilation (SIMV) modes. These modes allow both mandatory ventilation (including VCV, PCV and PRVC) and spontaneous ventilation (including spontaneous ventilation and pressure support ventilation - PSV).
- Spontaneous/ Continuous Positive Airway Pressure ventilation (SPONT/CPAP) mode. This mode only allows spontaneous ventilation.
- Bi Level ventilation (BIVENT) mode. This mode allows both mandatory ventilation (switching between high level CPAP and low level CPAP) and spontaneous ventilation.

NIV modes, include:

- NIV/CPAP mode – CPAP mode in NIV.
- NIV-T mode – A/C (PCV) mode in NIV.
- NIV-S/T mode – SPONT mode in NIV.

10.1.1 Assist/Control Ventilation

In Assist/Control ventilation, breaths are controlled by the ventilator (mandatory) or triggered by the patient (spontaneous) or triggered by the operator. When controlled by the ventilator, Ventilator Initiated Mandatory (VIM) breaths are flow controlled and time cycled, thus delivering an operator set volume (Tidal Volume) or pressure. Extra breaths, called Patient Initiated Mandatory (PIM) breaths, shall be possible if the patient overcomes the pre-set trigger level. PIM breaths are either pressure or flow triggered. If the trigger setting is adjusted so that the patient cannot trigger the ventilator, all breaths shall be VIM breaths, including operator triggered breaths.

1. Volume Control Ventilation:
In VCV, VIM breaths deliver the set volume (Tidal Volume - Vt).

Figure 10-1

Figure 10-2
2. PCV Pressure Controlled ventilation

In PCV, VIM breaths deliver the set Pressure (Pinsp) using a decelerating flow pattern.

3. Pressure Regulated Volume Control Ventilation

PRVC breaths will be delivered at a set rate and set volume (VT). The flow pattern resembles PCV. Inspiratory pressure will be regulated in PRVC to achieve the operator set volume (Tidal Volume). The first ventilation in PRVC will have a square flow waveform and each successive ventilation will have a ramp flow (decelerating) waveform.
PRVC ventilation operation shall be as follows:

When PRVC is selected, a volume controlled test breath, to the set tidal volume, will be delivered to the patient. The ventilator will set the target pressure for the first pressure control breath to the end inspiratory pressure of the test breath.

The next ventilation and all subsequent breaths will be delivered as pressure control breaths. The inspiratory pressure will be based on the dynamic compliance of the previous ventilation and the set tidal volume. Inspiratory pressure will be adjusted automatically by the ventilator to maintain the target tidal volume. The maximum step change between two consecutive breaths shall be 3 centimeters of water pressure. The maximum tidal volume delivered in a single ventilation shall be 1.5 times the Vt setting.

The test ventilation sequence shall be initiated when any of the following events occur:
Entering the Mode (PRVC) mode; Changing the set tidal volume while in PRVC; Delivered tidal volume >= 1.5 times the set volume; Flow termination of the test ventilation; activation of any of the following alarms - High Peak Pressure Alarm; Low Peak Pressure Alarm; Low PEEP alarm; Patient Circuit Disconnect Alarm; I-Time Limit; I:E Limit.

10.1.2 Synchronized Intermittent Mandatory Ventilation

SIMV is a ventilation mode where the patient is allowed to breathe spontaneously and the machine delivers VC mandatory breaths in synchrony with the patient's effort at the operator set rate and volume (or pressure). This is accomplished by a combination of spontaneous and mandatory windows that open and close. The type of ventilation delivered depends upon
whether the event during the window is patient initiated, operator initiated or time initiated. Spontaneous breaths occurring between mandatory breaths can be pressure supported. Synchronized breaths shall be either pressure or flow triggered. Back-up (Apnea) ventilation shall be provided when there is no patient trigger, mandatory ventilation or manual ventilation for a period that exceeds the apnea alarm setting. Backup Ventilation shall be available in SIMV - See section on Backup Ventilation for details on these modes and settings.

**Figure 10-5**

1. **SIMV (VCV)**
   
   In SIMV (VCV), mandatory ventilator breaths are volume controlled. Between the mandatory breaths the patient can breathe spontaneously. These spontaneous breaths can be pressure supported. Backup ventilation will be PCV.

2. **SIMV (PCV)**
   
   In SIMV (PCV), mandatory ventilator breaths are pressure controlled. Between the mandatory breaths the patient can breathe spontaneously. These spontaneous breaths can be pressure supported. Backup ventilation will be PCV.

3. **SIMV (PRVC)**
   
   In SIMV (PRVC), mandatory ventilator breaths are pressure-regulated volume controlled. Between the mandatory breaths the patient can breathe spontaneously. These spontaneous breaths can be pressure supported. Backup ventilation will be PCV.

### 10.1.3 Back-up Ventilation (Apnea Ventilation)

1. **Transition into Back-up Ventilation**

   The current Ventilation mode shall transition to Back-up Ventilation - PCV when an apnea occurs in SPONT/CPAP mode.
The current Ventilation mode shall transition to Back-up Ventilation - PCV when an apnea occurs in BIVENT mode.

The current Ventilation mode shall transition to Back-up Ventilation - PCV when an apnea occurs in NIV-S/T mode.

The current Ventilation mode shall transition to Back-up Ventilation - PCV when an apnea occurs in an SIMV mode.

2. Characteristics of Back-up Ventilation

Apnea ventilation settings include Pinsp, Tinsp or I:E.

3. Exiting Back-up Ventilation

Two methods to exit Back-up Ventilation shall be supported: trigger by patient or reset by operator.

When in Back-up Ventilation and the patient triggers two consecutive breaths and also the Expiratory Vt >= 0.5 x Inspiratory Vt for those two breaths, then the ventilator shall return to the Ventilation mode and settings prior to the Apnea event.

When in Back-up Ventilation and the operator resets the Apnea Alarm and confirms the reset by pressing the Encoder Knob, then the ventilator will return to the Ventilation mode and settings prior to the Apnea event.

Mode transitions when exiting Back-up Ventilation shall occur only when an Expiratory has been completed.

10.1.4 Spontaneous/CPAP Ventilation

In SPONT/CPAP Ventilation, the patient breathes spontaneously at a pressure level determined by the PEEP setting. Spontaneous ventilation may also be assisted by the ventilator at an operator set level of inspiratory pressure (Pressure Support). Inspiration is initiated by the patient and terminated when the inspiratory flow falls below an operator set percentage of the peak flow during this ventilation. During SPONT/CPAP, the patient determines the respiratory rate, and the patient and ventilator determine the inspiratory time and tidal volume. Ventilation detection shall be either pressure or flow triggered. PCV back-up ventilation shall be provided when there is no patient trigger, for a period that exceeds the apnea alarm setting. (See Section on Back-up ventilation for details)
10.1.5 BiLevel Ventilation (BIVENT)

In BIVENT, breaths shall be controlled by the ventilator (mandatory). Pressure controlled breaths are provided by switching between a high and low airway pressure in an adjustable time sequence. Spontaneous breaths can be pressure supported at the high and low pressure levels. When the expiratory time (Tlow) is less than the inspiratory time (Thigh) the displayed ventilation mode shall be Bi Vent APRV. PCV back-up ventilation shall be provided when there is no patient trigger, mandatory ventilation or manual ventilation for a period that exceeds the apnea alarm setting. (See Section on Back-up ventilation for details)
10.1.6 **Non Invasive/Continuous Positive Airway Pressure**

NIV/CPAP is a spontaneous mode of operation and no ventilator controlled breaths are provided. Throughout the ventilation cycle, an operator set pressure (CPAP) may be provided. In NIV/CPAP, the ventilator controls the airway pressure as the preset PEEP value.

Illustration shows NIV/CPAP. Pressure shall rise according to the selected rise time, with target pressure 1.5 cmH\textsubscript{2}O above PEEP to improve work of ventilating.
This illustration shows NIV/CPAP spontaneous breaths with Pressure Support.

Pressure support will be terminated when:
• Inspiratory flow returns to zero during phase 1 of inspiration, (i.e. when the patient exhales or fights the ventilator),
• Inspiratory flow in phase 2 of inspiration phase falls below a certain ratio (Esens) of the maximum value previously supplied when compared to the peak inspiratory flow supplied or
• Based on high Spont Inspiratory Time setting is exceeded

10.1.7 NIV-T

In NIV-T, breaths shall be controlled by the ventilator (mandatory) or shall be triggered by the patient (spontaneous) or shall be triggered by the operator. When controlled by the ventilator, breaths shall be pressure limited and time cycled, resulting in an operator set pressure (Pinsp) being delivered for an operator set period (Tinsp). Extra breaths shall be possible if the patient overcomes the pre-set trigger level or if the operator selects a manual ventilation. Patient triggered breaths shall be flow triggered.

10.1.8 NIV-S/T

NIV-S/T is a spontaneous mode of operation and no ventilator controlled breaths are provided. Throughout the ventilation cycle, spontaneous ventilation may also be assisted by the ventilator at an operator set of inspiratory pressure (Pressure Support). If the trigger time of the patient exceeds the setting value of the apnea time, the mode will enter into back-up ventilation.

NIV-S/Td is a spontaneous mode of operation and no ventilator controlled breaths are provided. Throughout the ventilation cycle, an operator set pressure (CPAP) may be provided. In NIV/CPAP, the ventilator controls the airway pressure as the preset PEEP value.
11 VG70 Ventilator System Specifications

⚠️ WARNING: Do not operate the Ventilator outside specified operating ranges or patient injury or equipment damage could result.

⚠️ NOTE: To add a measure of safety, an electronic hardware watchdog timer monitors the system operation at all times. In cases where the system software does not respond in a safe time period, the watchdog timer will reset the system. This enables the system software to be in control at all times.

11.1 System

11.1.1 General

This device complies with requirements of Medical Device Directive 93/42/EEC.

Standards

IEC 60601-1: The device classification is: Class I, Type B applied part (ventilator breathing tube and mask), type BF applied part (CO2 module), ordinary enclosed equipment without protection against ingress of liquids, continuous operation

ISO 80601-2-12

⚠️ WARNING: Equipment not suitable for use in the presence of a Flammable Anesthetic mixture with Air or with Oxygen or Nitrous Oxide.

Electromagnetic Compatibility (EMC)

According to IEC 60601-1-2

Patient Range (cm)

Adult: 60 – 260
Child: 30 – 140
NOTE: Leakage from VBS COMPLIANCE:

- 50 ml/min at 20 hPa for a VENTILATOR intended to provide DELIVERED VOLUME less than 50 ml;
- 100 ml/min at 40 hPa for a VENTILATOR intended to provide DELIVERED VOLUME between 300 ml and 50 ml;
- 200 ml/min at 50 hPa for a VENTILATOR intended to provide DELIVERED VOLUME greater than 300 ml.

### 11.1.2 Operating Conditions

Operating Temperature Range: +5 to +40°C  
Relative Humidity: 5 to 95% non-condensing  
Atmospheric Pressure: 700 to 1060 hPa

Enclosure Protection Rating IP21 per IEC 60529

### 11.1.3 Non-operating Conditions

Storage Temperature Range: -20 to +60°C  
Storage Relative Humidity: ≤ 95% non-condensing  
Storage Atmospheric Pressure: 700 to 1060 hPa

### 11.1.4 Power Supply

**Ventilator**

<table>
<thead>
<tr>
<th>Power Supply</th>
<th>100-240VAC 50/60 Hz</th>
</tr>
</thead>
<tbody>
<tr>
<td>DC Power</td>
<td>12-24 VDC</td>
</tr>
<tr>
<td><strong>Battery Backup (Standard)</strong></td>
<td>Two rechargeable lithium-ion battery modules, 14.4 V, 6.6 Ah each. Recharge time approximately 3.5 hours. Battery backup time of 120 minutes minimum with only standard internal battery.</td>
</tr>
<tr>
<td><strong>Maximum Power Consumption</strong></td>
<td>200VA</td>
</tr>
<tr>
<td><strong>AC Circuit Fuse</strong></td>
<td>UDA3.15</td>
</tr>
<tr>
<td><strong>DC Power Fuse</strong></td>
<td>GDA012</td>
</tr>
</tbody>
</table>
### Cart

<table>
<thead>
<tr>
<th>Output</th>
<th>Humidifier</th>
<th>Socket</th>
<th>Ventilator</th>
<th>Fuse</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>115V 2A 60Hz</td>
<td>115V 5A 60Hz</td>
<td>250V H3.5A</td>
</tr>
<tr>
<td></td>
<td></td>
<td>220V 1A 50Hz</td>
<td>220V 5A 50Hz</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>230V 1A 50/60Hz</td>
<td>230V 5A 50/60Hz</td>
<td></td>
</tr>
<tr>
<td>Input</td>
<td></td>
<td>115V 8A 60Hz</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>220V 8A 50Hz</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>230V 8A 50/60Hz</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

⚠️ **CAUTION:** DC power conforms to IEC 60601-1.

⚠️ **CAUTION:** The Power Supply should meet the above specifications.

⚠️ **WARNING:** Connecting any equipment that has not been supplied as a part of the ventilator system to the socket of the cart may generate the risk.

### 11.2 Ventilator

#### 11.2.1 General

<table>
<thead>
<tr>
<th>Dimensions</th>
<th>User Interface: 350 wide x 55 deep x 244 high (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ventilation Delivery Unit: 322 wide x 375 deep x 366 high (mm)</td>
</tr>
<tr>
<td></td>
<td>System with Cart (optional): 547 wide x 675 deep x 950 high (mm)</td>
</tr>
<tr>
<td>Weight (Approximate)</td>
<td>Total: 40 kg</td>
</tr>
<tr>
<td></td>
<td>User Interface: 2.5 kg</td>
</tr>
<tr>
<td></td>
<td>Ventilation Delivery Unit: 12.5 kg</td>
</tr>
<tr>
<td></td>
<td>Cart: 25 kg</td>
</tr>
<tr>
<td>Trigger Method</td>
<td>Flow and Pressure</td>
</tr>
<tr>
<td>Maximum limited pressure</td>
<td>80 cmH₂O</td>
</tr>
<tr>
<td>Maximum working pressure</td>
<td>80cmH₂O</td>
</tr>
<tr>
<td>Warm-up time</td>
<td>≥20 min</td>
</tr>
<tr>
<td>Pressure compensation</td>
<td>The ventilator has the automatic atmospheric pressure compensatory function.</td>
</tr>
</tbody>
</table>
11.2.2 Gas Supply

Supplied gases must be free of water, oil and particles.

Inlet Gas Pressure

$O_2$: 280 kPa to 600 kPa (41 – 87 psi)

Connection Standards Available

DISS, NIST

The maximum 10 s average input flow required by ventilator for $O_2$ is 60.69 LPM at a pressure of 280 kPa. For a 3 second average and at a pressure of 280kPa, the maximum averaged transient input flow required by the Ventilator is 64.44 LPM for $O_2$.

⚠️ **CAUTION:** The ventilator is a high flow device and should only be connected to a pipeline connection designed using a diversity factor that allows for the indicated high flow at a specified number of terminal outlets, in order to avoid exceeding the pipeline design flow, thereby minimizing the risk that the ventilator interferes with the operation of adjacent equipment.

11.2.3 Patient System Connectors

Male 22 mm Conical Fittings in accordance with ISO 5365-1.

11.2.4 User Interface

Attach to the ventilation delivery unit, the head mast, or other mounting system.

11.2.5 Acoustic Energy

The A-weighted sound pressure level of the ventilator is 45 dB; the A-weighted sound power level is 52 dB.

11.3 Standard Conditions Specifications

Error ranges in this document assume the following standard conditions:

- Ambient pressure: 101.3 kPa (1 atmosphere)
- Room temperature: 20°C
- Dry gases in patient system
- Inlet pressure: 345 kPa (50 psi)
11.4 Inspiratory Channel

Pressure Drop
Maximum 5 cmH₂O at a flow of 60 liters/min.

Rated Inspiratory Gas Pathway Resistance
Less or equal to 50 cmH₂O/L/s

Compliance
Maximum 2 mL/cmH₂O (With Fisher & Paykel MR 810 Humidifier and Patient Circuit of reusable silica gel).

Gas Delivery System
Microprocessor controlled valves.

Gas Delivery Device
Flow Range:
Adult: 1 to 180 liters/min
Child: 0 to 60 liters/min

Maximum Pressure Setting: 70 cmH₂O

NIV Max leakage compensation level
Adult: 60 lpm
Child: 30 lpm

11.5 Expiratory Channel

Pressure Drop (Resistance)
Maximum 5 cmH₂O at a flow of 60 lpm.

Rated Expiratory Gas Pathway Resistance
Less or equal to 20 cmH₂O/L/s

Compliance
Maximum 2 mL/cmH₂O (With Fisher & Paykel MR 810 Humidifier and Patient Circuit of reusable silica gel).

PEEP Regulation
Microprocessor controlled valve

PEEP Setting Range
Expiratory Flow Measurements

Range: 0 ~ 180 lpm.

Because of power failure or partial loss power, the ventilatory capacity is unnormal. When the ventilator supplies 60L/min volume, for the expiratory valve the pressure drop is 0.18kPa in inspiration, the pressure drop is 0.11kPa in expiration.

11.6 Monitoring

Inspiratory and Expiratory Minute Volume

Range: 0 ~ 60 lpm.
Accuracy: +/- 1 LPM or +/- 15% of measured value (whichever is greater)
Resolution: 0.1 lpm > 1 lpm, 0.001 lpm < 1 lpm

Inspiratory and Expiratory Tidal Volume

Range: 0 ~ 4000 mL
Adult Accuracy: +/- 25 ml or ±15% of the measured value (whichever is greater)
Child Accuracy: +/- 10 ml or ±10% of the measured value (whichever is greater)
Resolution: 1 mL

O₂ Concentration

Range: 18 ~100%
Accuracy: +/- 3 vol. %
Resolution: 1%

Airway Pressure

Range: -20 to 80 cmH₂O
Accuracy: ± (2 cmH₂O + 4% of reading)
Resolution: 1 cmH₂O

Measurement uncertainty

Volume: +/- 2% of reading or +/- 20 mL (whichever is greater)
Pressure: +/- 0.75% of reading or +/- 0.1 cmH₂O (whichever is greater)
O₂: +/- 1%

Airway Pressure and Flow Rate Waveform

Filtering of 5 samples is performed for smoothing.
11.7 Alarms

11.7.1 Allowed Alarm Settings

**Airway Pressure (upper limit)**
5 to 80 cmH₂O

**High Continuous Pressure**
Set PEEP level + 15 cmH₂O for at least 15 sec

**O₂ Concentration**
Set value +/- 6 vol % or <= 18 vol %

**Expired Minute Volume (Upper alarm limit)**
1 ~ 60 liters/min; OFF

**Expired Minute Volume (Lower alarm limit)**
0.1 ~ 40 liters/min; OFF

**Apnea Time**
10 ~ 60 sec; OFF

**Respiratory Frequency**
10 ~ 80 bpm; OFF

**Low End Expiratory Pressure**
1 ~ 35 cmH₂O; OFF
Or 1 ~ 20 cmH₂O; OFF (NIV modes only)

**End-Tidal CO₂ (Upper alarm limit)**
0.1% to 13.3% or 1 mmHg to 100 mmHg or 0.1 kPa to 13.3 kPa

**End-Tidal CO₂ (Lower alarm limit)**
OFF, 0.1% to 13.2% or OFF, 1 mmHg to 99 mmHg or OFF, 0.1 kPa to 13.2 kPa

11.7.2 Alarms Miscellaneous

**Gas Supply**
< 160 kPa (29 psi) +10% for 5 seconds or more
Alarm Silence/reset

Press this key to silence alarms for two minutes. This key also resets latched alarms.

Alarm Sound Pressure

The alarm sound pressure is above 60 dB at lowest volume setting at a distance of 1 meter from the front of the ventilator.

11.8 Ventilation Modes

11.8.1 Controlled Ventilation

Pressure Control (PCV)
Pressure controlled ventilation

Volume Control (VCV)
Volume controlled ventilation

Pressure Regulated Volume Control (PRVC)
Pressure regulated volume controlled ventilation

Noninvasive Pressure Control (NIV-T)
Noninvasive pressure controlled ventilation

11.8.2 Supported Ventilation

SPONT/CPAP+PSV
Spontaneous continuous positive airway pressure ventilation with pressure supported ventilation

NIV-S/T
Noninvasive pressure supported ventilation

11.8.3 Combined Ventilation

SIMV (PCV) + PSV
Synchronized intermittent mandatory ventilation based on pressure controlled ventilation with pressure supported ventilation
SIMV (VCV) + PSV

Synchronized intermittent mandatory ventilation based on volume controlled ventilation with pressure supported ventilation

SIMV (PRVC) + PSV

Synchronized intermittent mandatory ventilation based on pressure regulated volume controlled ventilation with pressure supported ventilation at high and low pressure levels

BIVENT

Pressure controlled ventilation that allows the patient the opportunity of unrestricted spontaneous ventilating with pressure support at high and low pressure levels

11.9 Trend Function

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Symbol</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peak Airway Pressure</td>
<td>$P_{peak}$</td>
</tr>
<tr>
<td>Plateau Airway Pressure</td>
<td>$P_{plat}$</td>
</tr>
<tr>
<td>Mean Airway Pressure</td>
<td>$P_{mean}$</td>
</tr>
<tr>
<td>End Expiratory Pressure</td>
<td>$PEEP$</td>
</tr>
<tr>
<td>Minimum Airway Pressure</td>
<td>$P_{min}$</td>
</tr>
<tr>
<td>Inspiratory Tidal Volume</td>
<td>$V_{ti}$</td>
</tr>
<tr>
<td>Expiratory Tidal Volume</td>
<td>$V_{te}$</td>
</tr>
<tr>
<td>Expiratory Minute Volume</td>
<td>$M_{Ve}$</td>
</tr>
<tr>
<td>Spontaneous Minute Volume</td>
<td>$M_{Vespont}$</td>
</tr>
<tr>
<td>Inspiratory/Expiratory Ratio</td>
<td>I:E</td>
</tr>
<tr>
<td>Total Ventilating Frequency</td>
<td>$f_{total}$</td>
</tr>
<tr>
<td>Spontaneous Ventilating Frequency</td>
<td>$f_{spont}$</td>
</tr>
<tr>
<td>Oxygen Concentration</td>
<td>$O_2$</td>
</tr>
<tr>
<td>Expiratory Resistance</td>
<td>$R_{exp}$</td>
</tr>
<tr>
<td>Dynamic Compliance</td>
<td>$C_{dyn}$</td>
</tr>
<tr>
<td>Rapid - shallow ventilating index</td>
<td>$RSBI$</td>
</tr>
<tr>
<td>Work of Ventilating</td>
<td>$WOB$</td>
</tr>
<tr>
<td>Leak</td>
<td>Leak NIV</td>
</tr>
</tbody>
</table>
## 11.10 Log Function

### Alarm/Event Log

Alarms
Ventilator Settings
Shortcut key functions

### Service Page

Technical alarms
Test results
Calibration results
Configuration log

## 11.11 Shortcut Key Functions

### Insp. Hold

Inspiratory Hold

### Exp. Hold

Expiratory Hold

### Nebulizer

Start and stop the Nebulizer operation

### Manual

Initiation of 1 ventilation in all ventilation modes except SPONT/CPAP

### Suction

Initiate and terminate the suction support process.
Disconnection detection – Automatic
Reconnection detection – Automatic
Pre-oxygenation – Max.3 minutes
Active Suction phase – Max.2 minutes
Post-oxygenation – 2 minutes

### Freeze

Freeze the current waveforms and loops, or unfreeze the Waveform display

### Screen Lock

Lock or unlock the touch screen
Main Menu

Return to the main menu

11.12 Communication/Interface

11.12.1 Nurse Call Port

The Ventilator has a modular jack configured to interface with external systems and is wired for normally open (N.O., close on alarm) signals. Contacts close on High priority alarms, loss of AC power and Speaker failure.

Floating DC contact
Voltage: Max. 50V
Current: Max. 200mA

Pin assignment:

1  Normally open  2  Normally open  3  Normally open
4  Common pin  5  Common pin  6  Common pin

11.12.2 Ethernet Port

Ethernet Port per IEE 802.3 to enable the VG70 to connect to external equipment such as an electronic health record.

⚠️ CAUTION:
• connection of the VG70 to an equipment could result in previously unidentified risks to patients, operators or third parties;
• the facility should identify, analyze, evaluate and control these risks
• subsequent changes to the Ethernet port could introduce new risks and require additional analysis; and
• changes to the Ethernet Port include:
  changes in Ethernet Port configuration;
  connection of additional items to the Ethernet Port;
  disconnecting items from the Ethernet Port;
  update of equipment connected to the Ethernet Port;
  update of equipment connected to the Ethernet Port.
11.12.3 RS-232 Port

Laptop-ready connection for user and service use in future software release.

11.12.4 Nebulizer Output Port

Output port for connection to nebulizer locates in the front of the ventilator. The whole nebulizer lasts for 30 minutes.

11.13 Accessories

Mobile Cart

Weight: 25Kg

Dimensions: 547 wide x 675 deep x 950 high (mm)

Gas Cylinder Kit

Capacity is two cylinders max, US E cylinders

Humidifier

Fisher & Paykel MR850, MR810

Breath circuit

Fisher & Paykel reusable breathing circuit:

| MV1006 (F&P) | 900MR761 Adult Circuit Kit |
| MV1005 (F&P) | 900MR742 Adult Single Limb Heated Circuit Kit |
| MV1007 (F&P) | 900MR780 Adult Circuit Kit, Dual Heated 4 LPM |
| MV1008 (F&P) | 900MR781 Infant Circuit Kit, Dual Heated 4 LPM |

Other recommended breathing circuit:

| 122000761 | Adult breathing circuit (Disposable): mask, headband, elbow, breathing circuit etc. Produced by INSPIRED MEDICAL |
| 122005487 | Adult Breathing Circuit (silicone)-mask, headband, elbow, breathing circuit etc. Produced by Liming Rubber |
| 122005490 | Pediatric Breathing Circuit-(silicone)-mask, headband, elbow, breathing circuit etc. Produced by Liming Rubber |
| 51005800 | Adult breathing circuit (Disposable). Produced by INSPIRED MEDICAL |
| 51008400 | Pediatric breathing circuit (Disposable). Produced by INSPIRED MEDICAL |
**Oxygen Sensor**: To be replaced every 2 years or as necessary. Part number: 210001975

**CO₂ Module**

<table>
<thead>
<tr>
<th>Part Number</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>240000415</td>
<td>IRMA CO₂ module</td>
</tr>
<tr>
<td>240000416</td>
<td>IRMA Adult airway adapter</td>
</tr>
<tr>
<td>240000417</td>
<td>IRMA Pediatric airway Adapter</td>
</tr>
</tbody>
</table>

### 11.14 Ventilating Parameters: Default Values and Allowed Settings (Standard Configuration)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Factory Set Default</th>
<th>Setting Range</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Child</td>
<td>Adult</td>
</tr>
<tr>
<td>ATC Tube Compensation</td>
<td>OFF</td>
<td>OFF</td>
</tr>
<tr>
<td>ATC Tube Type</td>
<td>ET</td>
<td>ET</td>
</tr>
<tr>
<td>ATC Tube Compensation (%)</td>
<td>80</td>
<td>80</td>
</tr>
<tr>
<td>ATC Tube Diameter (mm)</td>
<td>5.0</td>
<td>7.5</td>
</tr>
<tr>
<td>Backup Ventilation Pressure above PEEP (cmH₂O)</td>
<td>10</td>
<td>20</td>
</tr>
<tr>
<td>Backup Ventilation Tinsp (s)</td>
<td>0.6</td>
<td>1.0</td>
</tr>
<tr>
<td>Compliance Compensation</td>
<td>ON</td>
<td>ON</td>
</tr>
<tr>
<td>CPAP (cmH₂O) in NIV</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>Flow Trigger – Vsens (l/min)</td>
<td>2.0</td>
<td>2.0</td>
</tr>
<tr>
<td>Frequency - VCV bpm</td>
<td>30</td>
<td>15</td>
</tr>
<tr>
<td>Patient Height (cm)</td>
<td>100</td>
<td>150</td>
</tr>
<tr>
<td>I:E ratio</td>
<td>1:2</td>
<td>1:2</td>
</tr>
<tr>
<td>Maximum inspiratory</td>
<td>60</td>
<td>180</td>
</tr>
<tr>
<td>Parameter</td>
<td>Factory Set Default</td>
<td>Setting Range</td>
</tr>
<tr>
<td>-----------------------------------------------</td>
<td>---------------------</td>
<td>------------------------</td>
</tr>
<tr>
<td></td>
<td>Child</td>
<td>Adult</td>
</tr>
<tr>
<td>flow (liters/min)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maximum permitted pressure – safety valve</td>
<td>110</td>
<td>110</td>
</tr>
<tr>
<td>(cmH₂O)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mode (in NIV)</td>
<td>NIV/CPAP</td>
<td>NIV/CPAP</td>
</tr>
<tr>
<td>Mode (in Invasive Ventilation)</td>
<td>PCV</td>
<td>PCV</td>
</tr>
<tr>
<td>Nebulizer</td>
<td>OFF</td>
<td>OFF</td>
</tr>
<tr>
<td>Nebulizer Time (minutes)</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>NIV Rate (bpm)</td>
<td>30</td>
<td>15</td>
</tr>
<tr>
<td>O₂ Concentration (%)</td>
<td>40</td>
<td>40</td>
</tr>
<tr>
<td>PEEP (cmH₂O)</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>PEEP in NIV (cmH₂O)</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>Phigh(cmH₂O)</td>
<td>15</td>
<td>15</td>
</tr>
<tr>
<td>Plow (cmH₂O)</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>Press trig sensitivity level (cmH₂O)</td>
<td>-3</td>
<td>-3</td>
</tr>
<tr>
<td>Pressure level above PEEP (cmH₂O)</td>
<td>10</td>
<td>20</td>
</tr>
<tr>
<td>PS above PEEP (cmH₂O)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>PS – BIVENT above PEEP (cmH₂O)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>PS – NIV above PEEP (cmH₂O)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>SIMV rate (bpm)</td>
<td>20</td>
<td>4</td>
</tr>
<tr>
<td>Thigh(s)</td>
<td>0.6</td>
<td>1.0</td>
</tr>
<tr>
<td>Tinsp(s)</td>
<td>0.6</td>
<td>1.0</td>
</tr>
<tr>
<td>Tidal Volume (mL)</td>
<td>80</td>
<td>400</td>
</tr>
<tr>
<td>Tlow (s)</td>
<td>1.4</td>
<td>3.0</td>
</tr>
</tbody>
</table>
### Ventilator System Specifications

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Factory Set Default</th>
<th>Setting Range</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Child</td>
<td>Adult</td>
</tr>
<tr>
<td>Tpause (s)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Tslope(s)</td>
<td>0.1</td>
<td>0.1</td>
</tr>
</tbody>
</table>

#### Alarm Settings: Default Settings and Allowed Ranges

<table>
<thead>
<tr>
<th>Alarm Limits</th>
<th>Factory Set Default</th>
<th>Setting Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Airway Pressure, high limit (cmH₂O)</td>
<td>40</td>
<td>40</td>
</tr>
<tr>
<td>Airway Pressure, low limit (cmH₂O)</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>PEEP Low limit in Invasive Ventilation(cmH₂O)</td>
<td>OFF</td>
<td>OFF</td>
</tr>
<tr>
<td>PEEP Low limit in NIV (cmH₂O)</td>
<td>OFF</td>
<td>OFF</td>
</tr>
<tr>
<td>PEEP high limit in Invasive Ventilation (cmH₂O)</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>PEEP high limit in NIV (cmH₂O)</td>
<td>OFF</td>
<td>OFF</td>
</tr>
<tr>
<td>etCO₂ low limit %</td>
<td>4.0</td>
<td>4.0</td>
</tr>
<tr>
<td>etCO₂ low limit mmHg</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>etCO₂ low limit kPa</td>
<td>4.0</td>
<td>4.0</td>
</tr>
<tr>
<td>etCO₂ high limit %</td>
<td>6.5</td>
<td>6.5</td>
</tr>
<tr>
<td>etCO₂ high limit mmHg</td>
<td>49</td>
<td>49</td>
</tr>
<tr>
<td>etCO₂ high limit kPa</td>
<td>6.5</td>
<td>6.5</td>
</tr>
<tr>
<td>Expired Minute Volume, low limit (lpm)</td>
<td>0.5</td>
<td>1.0</td>
</tr>
<tr>
<td>Expired Minute Volume, high limit (lpm)</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>Expired Tidal Volume, low limit (mL)</td>
<td>50</td>
<td>250</td>
</tr>
<tr>
<td>Respiratory Frequency, Spontaneous, Upper limit (bpm)</td>
<td>OFF</td>
<td>OFF</td>
</tr>
<tr>
<td>Time, Apnea – until alarm (sec)</td>
<td>20</td>
<td>20</td>
</tr>
</tbody>
</table>
NOTE: Regarding occlusion detection and continuing pressure events:
Occlusion detection occurs in every ventilation mode. In the case that the Patient Airway Pressure is positive for more than 15 seconds at an unexpected level, the system will treat this continuing pressure event as an occlusion and behave accordingly.

NOTE: Regarding Nebulizer usage while on battery power:
The nebulizer function is disabled when the Ventilator is powered by either the internal or external battery.

Ventilating Parameters: Default Values and Allowed Settings (Standard Configuration).

11.15 Delivery Accuracy

Airway Pressure

+/- 10% of settings or +/- 3 cmH₂O (whichever is greater)

Tidal Volume

Adult Accuracy: +/- 10% of settings +/- 25 mL (whichever is greater)

Child Accuracy: +/- 10% of settings +/- 10 mL (whichever is greater)

O₂ Concentration

+/- 5% full scale for all modes

21% to 90% rising time (at worst case): 3 min 22 s

PEEP

+/- 10% of settings or +/- 2 cmH₂O (whichever is greater) for all modes

Measurement uncertainty

Volume: +/- 2% of reading or +/- 20 mL (whichever is greater)

Pressure: +/- 0.75% of reading or +/- 0.1 cmH₂O (whichever is greater)

O₂: +/- 1%
12 Pneumatic Diagram

Diagram showing a pneumatic system with various components such as filters, check valves, pressure switches, regulators, solenoid valves, mixing chambers, mufflers, turbines, proportional valves, oxygen sensors, pressure transducers, relief valves, and ambient valves. The system is designed to deliver oxygen to a patient through a nebulizer.
13 EMC Guidelines

Important information regarding Electro Magnetic Compatibility (EMC):

VG70 VENTILATOR needs special precautions regarding EMC and put into service according to the EMC information provided in the user manual; VG70 conforms to this IEC 60601-1-2:2014 standard for both immunity and emissions. Nevertheless, special precautions need to be observed.

- VG70 VENTILATOR with Following ESSENTIAL PERFORMANCE is intended used in Professional healthcare facility environment.

- WARNING: Use of VG70 VENTILATOR adjacent to or stacked with other equipment should be avoided because it could result in improper operation.

Below cables information are provided for EMC reference.

<table>
<thead>
<tr>
<th>Cable</th>
<th>Max. cable length, Shielded/unshielded</th>
<th>Number</th>
<th>Cable classification</th>
</tr>
</thead>
<tbody>
<tr>
<td>AC cable</td>
<td>5.0m Unshielded</td>
<td>1 Set</td>
<td>AC Power</td>
</tr>
</tbody>
</table>

- WARNING: The Use of accessories, transducers and cables other than those specified or provided by the manufacturer of the VG70 VENTILATOR could result in increased electromagnetic emissions or decreased electromagnetic immunity of this equipment and result in improper operation.

- WARNING: Portable RF communications equipment (including peripherals such as antenna cables and external antennas) should be used no closer than 30 cm (12 inches) to any part of the VG70 VENTILATOR, including cables specified by the manufacturer. Otherwise, degradation of the performance of this equipment could result.
EMI&EMS Compliance Table

### Table 1 - Emission

<table>
<thead>
<tr>
<th>Phenomenon</th>
<th>Compliance</th>
<th>Electromagnetic environment</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF emissions</td>
<td>CISPR 11 Group 1, Class A</td>
<td>Professional healthcare facility environment</td>
</tr>
<tr>
<td>Harmonic distortion</td>
<td>IEC 61000-3-2 Class A</td>
<td>Professional healthcare facility environment</td>
</tr>
<tr>
<td>Voltage fluctuations and flicker</td>
<td>IEC 61000-3-3 Compliance</td>
<td>Professional healthcare facility environment</td>
</tr>
</tbody>
</table>

**NOTE** The EMISSIONS characteristics of this equipment make it suitable for use in industrial areas and hospitals (CISPR 11 class A). If it is used in a residential environment (for which CISPR 11 class B is normally required) this equipment might not offer adequate protection to radio-frequency communication services. The user might need to take mitigation measures, such as relocating or re-orienting the equipment.

### Table 2 - Enclosure Port

<table>
<thead>
<tr>
<th>Phenomenon</th>
<th>Basic standard</th>
<th>EMC standard</th>
<th>Immunity test levels</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrostatic Discharge</td>
<td></td>
<td></td>
<td>±8 kV contact</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>±2kV, ±4kV, ±8kV, ±15kV air</td>
</tr>
<tr>
<td>Radiated RF EM field</td>
<td>IEC 61000-4-3</td>
<td></td>
<td>3V/m</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>80MHz-2.7GHz</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>80% AM at 1kHz</td>
</tr>
<tr>
<td>Proximity fields from RF wireless</td>
<td>IEC 61000-4-3</td>
<td></td>
<td>Refer to table 3</td>
</tr>
<tr>
<td>communications equipment</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rated power frequency magnetic fields</td>
<td>IEC 61000-4-8</td>
<td></td>
<td>30A/m</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>50Hz or 60Hz</td>
</tr>
</tbody>
</table>
## Table 3 – Proximity fields from RF wireless communications equipment

<table>
<thead>
<tr>
<th>Test frequency (MHz)</th>
<th>Band (MHz)</th>
<th>Immunity test levels</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Professional healthcare facility environment</td>
</tr>
<tr>
<td>385</td>
<td>380-390</td>
<td>Pulse modulation 18Hz, 27V/m</td>
</tr>
<tr>
<td>450</td>
<td>430-470</td>
<td>FM, ±5kHz deviation, 1kHz sine, 28V/m</td>
</tr>
<tr>
<td>710</td>
<td>704-787</td>
<td>Pulse modulation 217Hz, 9V/m</td>
</tr>
<tr>
<td>745</td>
<td>800-960</td>
<td>Pulse modulation 18Hz, 28V/m</td>
</tr>
<tr>
<td>780</td>
<td>1700-1990</td>
<td>Pulse modulation 217Hz, 28V/m</td>
</tr>
<tr>
<td>810</td>
<td>2400-2570</td>
<td>Pulse modulation 217Hz, 28V/m</td>
</tr>
<tr>
<td>870</td>
<td>5100-5800</td>
<td>Pulse modulation 217Hz, 9V/m</td>
</tr>
</tbody>
</table>
### Table 4 – Input a.c. power Port

<table>
<thead>
<tr>
<th>Phenomenon</th>
<th>Basic EMC standard</th>
<th>Immunity test levels</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Professional healthcare facility</strong></td>
<td></td>
<td><strong>Environment</strong></td>
</tr>
<tr>
<td>Electrical fast transients/burst</td>
<td>IEC 61000-4-4</td>
<td><strong>±2 kV</strong> 100kHz repetition frequency</td>
</tr>
<tr>
<td>Surges</td>
<td>IEC 61000-4-5</td>
<td><strong>±0.5 kV, ±1 kV</strong></td>
</tr>
<tr>
<td>Surges</td>
<td>IEC 61000-4-5</td>
<td><strong>±0.5 kV, ±1 kV, ±2 kV</strong></td>
</tr>
<tr>
<td>Conducted disturbances induced by RF fields</td>
<td>IEC 61000-4-6</td>
<td><strong>3V, 0.15MHz-80MHz</strong> 6V in ISM bands between 0.15MHz and 80MHz 80%AM at 1kHz</td>
</tr>
<tr>
<td>Voltage dips</td>
<td>IEC 61000-4-11</td>
<td><strong>0% Uₚ; 0.5 cycle</strong> At 05 cycle-11s between 0.15MHz and 80MHz zdstcommu</td>
</tr>
<tr>
<td>Voltage interruptions</td>
<td>IEC 61000-4-11</td>
<td><strong>0% Uₚ; 1 cycle</strong> and 70% Uₚ; 25/30 cycles Single phase: at 0t</td>
</tr>
</tbody>
</table>

### Table 5 – Signal input/output parts Port

<table>
<thead>
<tr>
<th>Phenomenon</th>
<th>Basic EMC standard</th>
<th>Immunity test levels</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Professional healthcare facility</strong></td>
<td></td>
<td><strong>Environment</strong></td>
</tr>
<tr>
<td>Electrostatic Discharge</td>
<td>IEC 61000-4-2</td>
<td><strong>±8 kV contact</strong> ±2kV, ±4kV, ±8kV, ±15kV air</td>
</tr>
<tr>
<td>Electrical fast transients/burst</td>
<td>IEC 61000-4-4</td>
<td><strong>±1 kV</strong> 100kHz repetition frequency</td>
</tr>
<tr>
<td>Conducted disturbances induced by RF fields</td>
<td>IEC 61000-4-6</td>
<td><strong>3V, 0.15MHz-80MHz</strong> 6V in ISM bands between 0.15MHz and 80MHz 80%AM at 1kHz</td>
</tr>
</tbody>
</table>
## ESSENTIAL PERFORMANCE:

<table>
<thead>
<tr>
<th>Requirement</th>
<th>Essential Performance</th>
</tr>
</thead>
</table>
| Oxygen level ALARM CONDITIONS| **O₂ Concentration Monitoring Range:** 18~100%  
**O₂ Concentration Setting Range:** 21~100%  
**High Oxygen Concentration Alarm:** Oxygen concentration exceeds the preset oxygen concentration +6% continuously for 30sec.  
**Low Oxygen Concentration Alarm:** Oxygen concentration is lower than the preset oxygen concentration -6% or lower than 18% continuously for 30sec  
**Oxygen Sensor Failure Alarm:** Checked during pre-inspection before use. With only 21% Air applied, O₂ reading after 20 seconds is ≤11% and ≥36% |
| AIRWAY PRESSURE              | **PEEP Setting Range:** 0-35 cmH₂O, 2 – 20 in NIV  
**Low PEEP Alarm:** PEEP measurement is lower than the alarm limit for three consecutive breath cycles.  
**High PEEP Alarm:** PEEP measurement is high than the alarm limit for three consecutive breath cycles  
**Occlusion Alarm:** Breathing circuit is occluded |
| Expired Volume               | **Expired volume range:** 20-2000ml  
**Expired volume monitoring range:** 0~4000 mL  
**High Expiratory Minute Volume:** Minute Volume exceeds the set limit, after 1min of ventilation, for three consecutive breath cycles or at most 10sec. The upper limit is larger than the lower limit by 1L at least  
**Low Expiratory Minute Volume:** Minute Volume is lower than the set limit within 1min after ventilation for three consecutive breath cycles or at most 10sec |
| Electrical supply failure    | **AC Failure:** During ventilator operation, when an AC power failure occurs and there is no battery power, the power board shall alarm for 120 seconds minimum. •When powered from batteries, an “AC Failure” alarm would occur. |
| INTERNAL ELECTRICAL POWER SOURCE near depletion | **Low Battery:** Under the battery operation, the remaining battery run time is less than 30min  
**Limited Battery Capacity:** Under the battery operation, the remaining battery run time is less than 10min |
| Gas supply failure           | **Low Airway Pressure Alarm:** Airway pressure is lower than the set limit for three consecutive breath cycles. |
| Gas failure cross flow       | **Low Oxygen Supply Pressure Alarm:** Oxygen supply is lower than 160kPa continuously for 5 sec. |