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The IPV® 1 may be covered by one or more patents.

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This manual was originally released and supplied in English. For a list of available translations, contact Customerservice.us@sentec.com. This manual may be revised or replaced at any time. Ensure this manual is the most current applicable version. To obtain the most recent version, contact Sentec Customerservice.us@sentec.com or visit Sentec.com.

⚠️ CAUTION: Federal law (U.S.) restricts this device to sale by or on the order of a physician.
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1: Introduction

The IPV® 1 system is intended for patients needing an airway clearance therapy for mobilization of secretions, lung expansion therapy, or the treatment and prevention of pulmonary atelectasis.

The IPV® 1 system is designed specifically for non-continuous institutional/hospital use to provide airway clearance and lung recruitment therapy. The system comprises a controller unit and breathing circuit called a Phasitron® 5. This circuit features a unique sliding venturi mechanism that provides a lung protective strategy while delivering high-frequency pulses, augmenting diffusive gas exchange throughout the lungs, promoting airway clearance and lung expansion.

About This Instruction Manual

This manual contains information for operating the IPV® 1 therapy system. Before operating the IPV® 1, the user must thoroughly read and understand these instructions for use.

It is the user’s responsibility to follow the instructions given and to keep the instructions for use near the device to ensure correct operation. If the safety instructions are not followed, the patient may be at risk.

This section contains the following:
- Related documents and additional resources
- Symbol definitions
- Safety information, including warnings and cautions
- Technical assistance information

Related Documents and Resources

The current version of this manual, specifications, clinical studies, and additional information is available at:

Customerservice.us@sentec.com
Glossary of Symbols

The table below summarizes symbols that may be used on the IPV® 1 (including all its related parts), on the packaging, and in the associated documentation. These symbols indicate information essential for proper use; the order of their appearance is not prioritized.

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>![Manufacturer symbol]</td>
<td>Manufacturer</td>
<td>Indicates the medical device manufacturer.</td>
</tr>
<tr>
<td>![Date of Manufacture symbol]</td>
<td>Date of Manufacture</td>
<td>Indicates the date when the medical device was manufactured.</td>
</tr>
<tr>
<td>![Country of Manufacture symbol]</td>
<td>Country of Manufacture</td>
<td>Identifies the country of manufacture of products. (CC to be replaced by country code.)</td>
</tr>
<tr>
<td>![Use-by date symbol]</td>
<td>Use-by date</td>
<td>Indicates the date after which the medical device is not to be used.</td>
</tr>
<tr>
<td>![Batch code symbol]</td>
<td>Batch code</td>
<td>Indicates the manufacturer’s batch code so that the batch or lot can be identified.</td>
</tr>
<tr>
<td>![Catalogue number symbol]</td>
<td>Catalogue number</td>
<td>Indicates the manufacturer’s catalogue number so that the medical device can be identified.</td>
</tr>
<tr>
<td>![Medical Device symbol]</td>
<td>Medical Device</td>
<td>Indicates that the device is a medical device.</td>
</tr>
<tr>
<td>![Serial number symbol]</td>
<td>Serial number</td>
<td>Indicates the manufacturer’s serial number so that a specific medical device can be identified.</td>
</tr>
<tr>
<td>![Fragile, handle with care symbol]</td>
<td>Fragile, handle with care</td>
<td>Indicates a medical device that can be broken or damaged if not handled carefully.</td>
</tr>
<tr>
<td>![Keep dry symbol]</td>
<td>Keep dry</td>
<td>Indicates a medical device that needs to be protected from moisture.</td>
</tr>
<tr>
<td>![Temperature limit symbol]</td>
<td>Temperature limit</td>
<td>Indicates the temperature limits to which the medical device can be safely exposed (upper and lower limits of temperature are indicated adjacent to the upper and lower horizontal lines).</td>
</tr>
<tr>
<td>![Humidity limitation symbol]</td>
<td>Humidity limitation</td>
<td>Indicates the range of humidity to which the medical device can be safely exposed (humidity limitation indicated adjacent to the upper and lower horizontal lines).</td>
</tr>
<tr>
<td>Symbol</td>
<td>Name</td>
<td>Description</td>
</tr>
<tr>
<td>--------</td>
<td>-------------------------------------------</td>
<td>------------------------------------------------------------------------------</td>
</tr>
<tr>
<td></td>
<td>Atmospheric pressure limitation</td>
<td>Indicates the atmospheric pressure limits to which the medical device can be safely exposed.</td>
</tr>
<tr>
<td></td>
<td>Single patient - multiple use</td>
<td>Indicates a medical device that may be used multiple times (multiple procedures) on a single patient.</td>
</tr>
<tr>
<td></td>
<td>Consult instructions for use</td>
<td>Indicates the need for the user to consult the instructions for use.</td>
</tr>
<tr>
<td></td>
<td>Mandatory action: refer to instruction manual</td>
<td>Indicates that the instruction manual must be read.</td>
</tr>
<tr>
<td></td>
<td>Caution Symbol</td>
<td>Indicates that caution is necessary when operating the device or that the current situation needs operator awareness or operator action to avoid undesirable circumstances.</td>
</tr>
<tr>
<td></td>
<td>Not Made with Natural Rubber Latex</td>
<td>Natural rubber latex was not used as a material in the manufacture of a medical product, its container and/or packaging.</td>
</tr>
<tr>
<td></td>
<td>Does Not Contain the Phthalate Plasticizers DEHP, DIBP, DBP, or BBP</td>
<td>Indicates product that does not contain the phthalate plasticizers DEHP, DIBP, DBP, or BBP.</td>
</tr>
<tr>
<td></td>
<td>Prescription only</td>
<td>Caution: Federal Law (U.S.) restricts this device to sale by or on the order of a physician.</td>
</tr>
<tr>
<td></td>
<td>Unique device identifier</td>
<td>Indicates a carrier that contains unique device identifier information.</td>
</tr>
</tbody>
</table>

**Safety Information**

This section contains important information for users. Please pay close attention to the **Warnings** and **Cautions**, and their associated consequences. Always exercise appropriate caution while using the IPV® 1 system.
A **WARNING** indicates the possibility of injury, death, or other serious adverse reaction associated with the use or misuse of the device.

<table>
<thead>
<tr>
<th>General Warnings</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>• Before use, read the instruction manual.</td>
<td></td>
</tr>
<tr>
<td>• The IPV® 1 system shall not be used as a life support device.</td>
<td></td>
</tr>
<tr>
<td>Supplemental oxygen must be prescribed for patients for whom it is indicated,</td>
<td></td>
</tr>
<tr>
<td>and the O2 saturations should be monitored.</td>
<td></td>
</tr>
<tr>
<td>• An IPV® treatment mobilizes secretions. Drainage techniques (such as controlled</td>
<td></td>
</tr>
<tr>
<td>coughing) are particularly important for patients with reduced ability to</td>
<td></td>
</tr>
<tr>
<td>spontaneously cough.</td>
<td></td>
</tr>
<tr>
<td>• Do not use with uncooperative patients.</td>
<td></td>
</tr>
<tr>
<td>• The intended user must continue to monitor the patient for progression of</td>
<td></td>
</tr>
<tr>
<td>symptoms and re-evaluate the indications for therapy.</td>
<td></td>
</tr>
<tr>
<td>• For neonates, use higher frequencies and carefully adjust amplitude from zero</td>
<td></td>
</tr>
<tr>
<td>to desired effect. Always monitor neonates and young children during treatment.</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Alteration/Assembly</th>
<th>Carefully follow assembly instructions outlined in this instruction manual. Assembly deviation/alteration could cause the IPV® 1 to malfunction. The IPV® 1 system cannot be used in an MRI environment.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Before Using</td>
<td>Before activating the IPV® 1, be sure to complete pre-checks to ensure proper operation.</td>
</tr>
<tr>
<td>Equipment</td>
<td></td>
</tr>
<tr>
<td>Breathing Circuits</td>
<td>Do not substitute any other breathing circuit for the Phasitron® 5 UC breathing circuit. Do not change the configuration of the breathing circuit. The Phasitron® 5 UC breathing circuit is specifically designed for use with the IPV® 1. Do not use any third-party circuits as their use has not been tested by Percussionaire; performance and safety cannot be guaranteed.</td>
</tr>
<tr>
<td>Patient Interface</td>
<td>Do not use a mouthpiece during IPV® 1 therapy for patients less than five years of age. Make sure mask is appropriately sized, covering mouth and nose but not the eyes, and not overlapping the chin.</td>
</tr>
<tr>
<td>Guidelines</td>
<td>Instructions in the manual are suggested guidelines for trained respiratory therapists and clinicians working under a supervising physician. Carefully select settings based on clinical judgment, the needs of the patient, and the benefits, limitations, and characteristics of IPV® 1. Always follow hospital or institutional protocols.</td>
</tr>
<tr>
<td>Infection Control</td>
<td>Dirty or contaminated equipment is a potential source of infection. Never attempt to reuse single-patient components or accessories between patients. The Phasitron® 5 UC breathing circuit is single patient. Replace the entire circuit when it cannot be clean “as new.”</td>
</tr>
<tr>
<td>Pre-Use Checks</td>
<td>Complete pre-use checks before starting therapy on a patient. Do not use the device if any problems are noted. Contact a qualified service technician. Failure to comply could cause injury or death to the patient.</td>
</tr>
<tr>
<td>Setup</td>
<td>Ensure the patient circuit is oriented correctly. Carefully follow setup directions given in the instructions for use.</td>
</tr>
<tr>
<td>Residual Risk/</td>
<td>The patient must notify the physician and the RN (Registered Nurse) or the RRT (Registered Respiratory Therapist) if they experience increased shortness of breath; significant changes in heart rate or rhythm, blood pressure, or skin color; marked diaphoresis; fatigue; or emesis.</td>
</tr>
<tr>
<td>Information to the Patient</td>
<td></td>
</tr>
</tbody>
</table>
A CAUTION indicates the possibility of a problem with the device associated with its use or misuse, such as a device malfunction, device failure, device damage, or other property damage.

### Cautions/Precautions
- All persons providing IPV® treatment must be trained in the use of Percussionaire devices.
- Therapists should evaluate how their patients tolerate the treatment. Auscultation and observation of the mechanical vibrations of the chest and abdomen are primary indicators of effective treatment.
- Care should be taken to appropriately suction secretions as they mobilize into the upper airways.
- Use only genuine Percussionaire parts and accessories.

### Cleaning and Disinfection
**IPV® 1 Driver:**
- Clean as required. Do not spray any liquid directly onto the device.
- Only use non-abrasive cotton cloths, cleaning wipes, and paper towels.
- Use any cleaning and disinfection solutions and products with caution.
- Use only the cleaning and disinfecting procedure recommended by the manufacturer.

**Breathing Circuit:**
- Disconnect tubing from Phasitron® 5 UC before cleaning.
- Do not wash or submerge the antibacterial filter or tubing harness. Use a clean, damp cloth to wipe the exterior of the tubing harness.
- Use of cleaning methods not outlined in these instructions for use may damage parts of the Phasitron® 5 UC breathing circuit kit.

### Clinician Training
All persons providing IPV® therapy must be trained in the use of the IPV® 1, the functions, and the settings.

All persons operating the IPV® 1 must read and understand the manual before using the device. All persons operating the IPV® 1 must read and understand the manual before using the device.

### Do NOT Cover Device
Do not cover the IPV® 1 during use. Do not place objects on top of the IPV® 1.

### Malfunctions
Do not use the device if any problems are noted. Contact qualified service technician.

### Maintenance
Maintenance is to be performed by authorized service technicians only. Send the device for maintenance and service following the recommendations provided in this manual.

### Incoming Air/Oxygen Pressure Settings
Always check the incoming air/oxygen pressure settings before connecting to the IPV® 1 device.

### Technical Assistance
For technical information and assistance, to request service, or to order parts, use one of the following methods of contact:

- Telephone: Technical service at 1-877-425-8746.
- Email: Customerservice.us@sentec.com
Indications for Use

The IPV® 1 system is used where Airway Clearance Therapy (ACT) is indicated. Airway Clearance Therapy is indicated for mobilization of secretions, lung expansion therapy, and the treatment and prevention of pulmonary atelectasis.

Intended Purpose

The IPV® 1 driver (control unit) is an active medical device intended to be used with adults, pediatrics, and neonates within the hospital/clinical/doctor’s office environment to provide Airway Clearance Therapy (ACT) by clinically trained professionals or respiratory technicians.

Intended Use Environment

The IPV® 1 system is intended to be used in the hospital and clinical environment or other institutions (doctor’s office) that have respiratory treatment facilities and accredited professionals to provide ACT.

Intended User Profile

The IPV® 1 system is intended to be used by healthcare professionals (including respiratory technicians), clinically trained in airway clearance therapy.

Indications for Use

The IPV® 1 system is used where Airway Clearance Therapy is indicated. Airway Clearance Therapy is indicated for mobilization of secretions, lung expansion therapy, and the treatment and prevention of pulmonary atelectasis.

Expected Clinical Benefits of IPV®

The IPV® 1 system is a form of IPV® used for ACT. IPV® is used in many respiratory conditions (including, among others, chronic obstructive pulmonary disease, cystic fibrosis, and Duchenne muscular dystrophy), where the following clinical and/or physiological benefits are desired:

- Lung recruitment
- Improved lung function
- Improved gas exchange
- Increased secretion mobilization
- Decreased work of breathing
- Decreased Length of Stay
- Reduced escalation of ventilatory support
The IPV® 1 system is indicated to provide IPV® as a form of airway clearance therapy. During operation, the system provides a continuous, dense aerosol mist to reduce the adhesive and cohesive forces of retained airway secretions.

### Contraindications

<table>
<thead>
<tr>
<th>Contraindication</th>
<th>Condition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Untreated tension pneumothorax</td>
<td>Untrained or unskilled operator</td>
</tr>
<tr>
<td>History of pneumothorax</td>
<td>Myocardial infarction</td>
</tr>
<tr>
<td>Pulmonary hemorrhage</td>
<td>Vomiting</td>
</tr>
<tr>
<td>Recent pneumonectomy (without functioning chest tube)</td>
<td>Pulmonary air leak</td>
</tr>
</tbody>
</table>
3: Principles of Operation

The IPV® 1 system is a device that provides IPV® as a form of airway clearance therapy. During operation, the system provides continuous dense aerosol mist to reduce the adhesive and cohesive forces of retained airway secretions. IPV® provides a “Pulsatile Flow” that mobilizes and helps remove mucus and debris, while gently recruiting alveoli and restoring FRC. The mechanism by which IPV® may promote pulmonary secretion clearance is its high-frequency percussive flow patterns that disrupt the adhesive and cohesive properties of mucus and other debris. The resulting exiting gas flow out of the tracheal bronchial tree helps to further transport the loosened mucus in a cranial direction.

Airway clearance therapy techniques all have a series of different physiological mechanisms used for “unblocking” the obstruction:

- Increase of expiratory flow
- Oscillation of the airflow
- Increase in the gas exchange

The aim of airway clearance therapy is to reduce airway obstruction caused by secretions occupying the airway lumen and so help to prevent respiratory tract infections, and to re-expand collapsed areas of the lung, thus supporting improved gas exchanges and decreased inflammatory response.
4: Description

IPV® 1 System

Front Panel

1. Digital Display
2. AMPLITUDE Adjustment Knob
3. GAS SOURCE Selector Knob
4. Breathing Circuit Connector
5. FREQUENCY Adjustment Knob
6. Phasitron® 5 UC Holder
### Control Functions

<table>
<thead>
<tr>
<th>Control</th>
<th>Functions</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. FREQUENCY</td>
<td>FREQUENCY determines the rate of high-frequency percussive pulses delivered to the patient.</td>
</tr>
<tr>
<td>2. AMPLITUDE</td>
<td>AMPLITUDE determines the pressure delivered to the patient.</td>
</tr>
<tr>
<td>3. GAS SOURCE</td>
<td>GAS SOURCE selects air or oxygen, or turns IPV® 1 off to stop therapy.</td>
</tr>
<tr>
<td>4. Digital Display</td>
<td>Digital Display reads and displays Mean Airway Pressure (MAP), Pulse Frequency Rate, Usage Time, and Pulse Amplitude.</td>
</tr>
</tbody>
</table>
The IPV® 1 connects to hospital single gas source or blended gas. Single or dual air/oxygen gas connections are standard.

⚠️ CAUTION: Bleed must remain unrestricted.
5: Digital Display

A digital display found on the front panel of the IPV® 1 device provides feedback of the patient proximal airway pressures, pulse frequency, and elapsed treatment time.

The display has six different operating modes: POST, Wake, Active, Report, Sleep, and Fault.

Note: The display will not awaken until feedback from proximal line on the Phasitron® 5 UC sees pressure.

POST (Power-On Self-Test)

The power-on self-test (POST) is a succession of built-in diagnostic tests performed by the digital display when batteries are installed.

When batteries are installed in the digital display, the software displays the System Information page for 15 seconds. This page includes the battery voltage, total usage time, software revision, and serial number.

This start-up mode allows the software to perform tests on the hardware that are part of POST. If any errors are detected in POST, the display enters Fault mode.

The POST check requires that the measurement port be left disconnected and exposed to the atmosphere for the entire duration.

NOTE: Do not install the digital display into the IPV® 1 until the POST check is complete and the screen is blank (indicating Sleep mode).
Wake Mode

To wake up the display, ensure the IPV® 1 pressure is greater than 7 cmH₂O/hPa at the Phasitron® patient-delivery port for more than 1 second.

The display remains on for the first 15 seconds, showing the bar-graph timer. If usage is stopped within 12 seconds, the display enters Report mode. After 15 seconds, the current session continues counting from 16 seconds, then turns into Active mode.

Active Mode

**Display Metrics:** Pulse Frequency Rate, Mean Airway Pressure, Usage Timer, Pulse Amplitude Pressure.

At 16 seconds, the display enters Active mode, a numeric display of Pulse Amplitude, Pulse Frequency, Mean Airway Pressure, and the Usage Timer.

- **Pulse Amplitude** - calculated from the pressure measurements at the moment of instantaneous peak and trough amplitude averaged over 5 seconds.

- **Pulse Frequency Rate** – the current measurement.

- **Mean Airway Pressure (MAP)** - averages Pulse Amplitude over 5 seconds. At 100 samples per second, this is an average of 500 measurements.

- **Usage Timer** – displayed in minutes and seconds, the Usage Timer is the total usage time of the current session. The timer can display a maximum of 59 minutes and 59 seconds. If usage has stopped for more than 5 minutes, the timer resets and starts over.

**NOTE:** To display most recent usage duration time, see Report mode next page.
Report Mode

The Session Timer and the Total Usage Timer (A) are displayed for 2 seconds, followed by the System Information Page (B) for 2 seconds, alternating. The alternating page display continues for 5 minutes, or until usage resumes and the display enters Active Mode.

During the 5-minute period, a horizontal bar graph indicates the time by moving from left to right at a fixed rate. After 5 minutes of no usage, the system information page is no longer displayed, and the time display (C) flashes, 2 seconds on, 2 seconds off, for an additional 25 minutes. The display enters Sleep mode after 25 minutes.

**NOTE:** When the IPV® 1 is turned off, the measurements will drop to zeros after a few seconds.

Sleep Mode

In Sleep mode, the display is off, but the microcontroller continues to sample and calculate the pressure at the measuring port 5 times a second. Over any 3-second period, if the pressure is greater than 7 cmH₂O/hPa at the Phasitron® 5 UC patient-delivery port for more than 1 second, the display enters Wake mode.
Fault Mode

In Fault mode, the display will show an error message on the display stating, “Contact Factory for Service.”

The displayed information includes the software revision, digital display serial number, the total usage time, and an error code for the exclusive use of the factory. (See System Failure Note below.)

NOTE: Pressure faults are triggered by a very high continuous pressure for more than 5 seconds during Wake and Active modes.

In all other modes, the software continuously monitors the hardware for errors and verifies that each data sample is valid. If an error is detected, the software logs the error and reboots the processor. Rebooting allows the display to recover from a transient error. After reboot, the processor returns to the same mode it was in before the reboot. If more than one error is detected in any 10-second period, it is considered “fatal,” and the software enters Fault mode.

NOTE: If System Failure screen is displayed, remove batteries for 30 seconds. Replace batteries (note that positive terminals face the same direction), and wait 30 seconds until the screen turns off. If POST runs correctly, the display may be used. If System Failure screen recurs, contact customer support.

Fault Detection

The display has both hardware and software fault detection. The dedicated hardware “watchdog” runs on an independent clock source and can continue to operate even if the main microprocessor’s clock fails, or the microcontroller pauses in any way. The independent fault detection is reset each time a valid pressure reading (free of hardware and software errors), is obtained. The software fault detection watchdog detects when a software task fails to complete within a specified time, logs the error, and resets the processor.

Fault Logging

The software keeps track of several types of hardware and data faults specific to the display; it does not record patient-related data. All faults are logged in the microcontroller’s memory and are retained even if the batteries are removed. If multiple faults happen within 10 seconds of each other, the display stops normal operation and enters Fault mode. In this mode, a subset of the collected fault information is shown on the display. This data is intended for service use only.

The user can exit the Fault mode by removing and replacing the batteries. This resumes normal operation of the IPV® 1 display but does not erase the faults stored in memory or fix the problem that caused the fault.
When setting up the IPV® 1 device for the first time, the digital display must be removed to access the battery-disconnect tabs for removal.

**NOTE:** When the display powers on for the first time, it will ensure the correct atmospheric pressure calibration at startup.

1. Press in on the display’s bezel and twist counterclockwise (left) approximately 20 degrees.
2. Gently pull on the display’s bezel to remove it from the IPV® 1 housing.
3. Remove the two battery-disconnect tabs to insert the batteries.
4. When the screen goes blank, the display is ready to be reinstalled into the IPV® 1.

**NOTE:** Refer to display instructions POST mode (page 12) to verify display operation.

---

**Digital Display - Setup**

![Side view of display module](image)

**Install batteries into display**

**USB serial port only used for firmware upgrades**

**Pull up to remove both tabs**

**Installing batteries: note positive terminals face same direction.**
6: Setup

Roll Stand Assembly

The IPV® 1 mounts onto a standard 0.75- to 1.5-inch (19 mm to 38 mm) hospital pole stand.
Attaching IPV® 1 to Stand

To mount device onto pole, open clamp.

Use adjuster nut to set pole thickness.

Close and lock clamp.
The patented Phasitron® 5 uses a unique venturi as a “clutch” mechanism to protect the lung from overpressure. By automatically adjusting to the resistance of the lung, the Phasitron® 5 precisely and safely delivers the optimal amount and flow of air required by the alveolar space. When lung resistance is low, as in a compliant lung, all the pulsed air from the IPV® 1 enters the mouth of the venturi tube. This low-pressure entrained air automatically fills the available space in the lung. The Phasitron® 5 continuously and instantaneously adjusts to keep a gentle and safe air pressure, even in a compromised lung.

Only use Percussionaire accessories designed specifically for use with the IPV® 1 device. Function is not guaranteed with any third-party equipment.
Breathing Circuit - Phasitron® 5 UC - Components

Breathing Circuit - Phasitron® 5 UC - Configurations

The Phasitron® 5 UC breathing circuit can be used with a mouthpiece or face mask. It can also be used without a mouthpiece or mask.

NOTE: It is recommended to use a resuscitation mask or similar non-vented mask for all patients under five years of age. International guidelines recommend that an appropriately sized face mask should cover the mouth and nose, but not the eyes, and should not overlap the chin. Careful consideration must be given to mask size, fit, and hold; a good seal with minimized leak is important for establishing effective therapy.

WARNING: Due to the requirements for physiological and neurological development, using a mouthpiece during therapy is not recommended for patients less than five years of age.
The Phasitron® 5 UC breathing circuit connects into the bottom of the IPV® 1 device.

Connecting Tubing to Phasitron® 5

Connect yellow tubing quick-connect fitting to nebulizer bowl.

**NOTE:** The tubing connectors will only fit onto the correct part

Press red tubing onto conical connector at rear of Phasitron® 5 body.

Connect the clear tubing quick-connect fitting to the cap at the rear of the Phasitron® 5 body.
Adding Liquid Solution

Twist nebulizer bowl to the left to open. Add liquid solution, e.g., saline or other diluent as directed by physician.

Apply the lid and twist the nebulizer bowl to the right to close.
7: Pre-Use Check

1. Verify the **GAS SOURCE** knob in the “OFF” position.

2. Connect the IPV® 1 to medical 50 psi oxygen or air gas supply.

3. Connect a Siemens®-style 1-liter ventilator test lung (A) to the Phasitron® 5 (B).

4. Connect the Phasitron® 5’s Unified Connector (C) to the IPV® 1 (D).

5. Rotate **GAS SOURCE** knob to the AIR or O₂ setting, depending on connection made in step #2.

6. Rotate the **AMPLITUDE** control knob fully counterclockwise.
7. Rotate the **FREQUENCY** control knob fully counterclockwise.

8. Confirm frequency rate of 300 or above.

**NOTE:** Displayed pressure and time are for illustrative purposes only.

9. Verify the mean airway pressure (MAP) indicated on the digital display is greater than 15 cmH₂O.

**NOTE:** Displayed pulse amplitude pressure, time, and frequency are for illustrative purposes only.

10. Rotate the **FREQUENCY** control knob fully clockwise.

11. Confirm frequency rate of 100 or below.

**NOTE:** Displayed pulse amplitude pressure, time, and mean airway pressure (MAP) are for illustrative purposes only.
12. Verify the mean airway pressure (MAP) indicated on the digital display is greater than 10 cmH₂O.

**NOTE:** Displayed pulse amplitude pressure, time, and frequency are for illustrative purposes only.

13. Rotate the **FREQUENCY** control knob to the center, straight up position.

14. Rotate the **AMPLITUDE** control knob fully clockwise.

15. Rotate the **GAS SOURCE** knob to the “OFF” position.

16. Disconnect gas source supply.
8: Prepare for Patient-Airway Connection

To prepare for patient-airway connection, complete the following steps:

1. Rotate **AMPLITUDE** control knob clockwise to stop (off).

2. Connect the IPV® 1 to 50 psi (3.4 bar) gas source. Ensure **GAS SOURCE** switch is switched to **OFF**.

3. Make sure patient is in an upright, comfortable position or lying with head and shoulders elevated by pillows.

**NOTE:** *Patient’s gravitational position is not a factor with IPV®.*

4. Auscultate patient for breath sounds, heart and respiratory rate, or follow institutional guidelines.

5. Place liquid solution into nebulizer bowl to a maximum of 20 cc. Use normal saline or other diluent as directed by physician.

6. Connect Phasitron® 5 UC breathing circuit kit to the IPV® 1 device.

7. Rotate the **FREQUENCY** control knob fully counterclockwise.

8. Connect Phasitron® 5 to the patient using mouthpiece, mask, or direct connection to patient airway.

9: Administer IPV® Therapy

1. Turn IPV® 1 on by rotating **GAS SOURCE** knob to relevant gas source. Slowly rotate **AMPLITUDE** control knob until a visible chest wiggle is observed.

2. When using a mouthpiece, instruct the patient to inhale and exhale through the pulses until a visible chest wiggle is observed throughout the whole chest or percussions can be auscultated in all lung fields.

**NOTE:** *Most patients will initially allow percussive bursts of air to leak through their lips or nose at the expense of an observable chest movement (wiggle).*

3. Start to notice the chest movement (wiggle), as the patient exhales through the mouthpiece. Advise the patient to relax, taking normal (spontaneous), breaths through the pulses whenever they desire.

4. Instruct the patient to keep their lips and cheeks tight. As the patient learns to prevent air from leaking out of the lip seal around the mouthpiece, the **FREQUENCY** control knob can be gradually rotated clockwise.

**NOTE:** *Some patients may require a nose clip to prevent venting from the nose, or an appropriate-size mask if unable to maintain a tight lip seal or venting through the nose.*
5. After the patient obtains the ability to prevent the leaking of percussive air deliveries from the nose and around the lips, the entire percussion rate range should be scanned by rotating the **FREQUENCY** control knob. This works to raise secretions from the bronchial airways.

**NOTE:** *Pay attention to patient comfort and make treatment adjustments accordingly.*

6. As the learning period progresses, the selected source pressure may be increased for effective endobronchial percussion by assessing chest percussion (wiggle).

7. Continue IPV® therapy for 15 to 20 minutes, or per orders.

8. When treatment is complete, turn the IPV® 1 off by rotating the **GAS SOURCE** knob to **OFF**. The Phasitron® 5 UC should be rinsed or cleaned and stored in a clean bag, or per hospital infection control policy, until the next treatment.

**NOTE:** The Phasitron® 5 UC breathing circuit kit is for SINGLE patient use only.

**NOTE:** *It is advised to follow manufacturer’s recommendation for cleaning (Chapter 10).*
Always clean the controller between patients and when visibly soiled. Use only approved cleaning wipes (CaviWipe®) to wipe the controller and stand to remove excess soil. Use additional CaviWipes if needed to remove visible soil. A visual inspection should be performed to ensure soil has been removed.

**NOTE:** Percussionaire® makes no claims regarding the efficacy of the listed chemicals or processes as a means for controlling infection. Consult your hospital’s infection control officer or epidemiologist. To clean or sterilize mounted devices or accessory equipment, refer to the manufacturers’ instructions for the devices or equipment.

⚠️ **CAUTION:** Do not spray any cleaning solution onto the controller or stand.

⚠️ **CAUTION:** Do not allow liquids to access the controller.

### Digital Display

1. Take care not to damage or scratch the digital display with fingernails, rings, or jewelry.
2. Do not apply pressure to the digital display.
3. Use CaviWipes if needed to remove visible soil.
4. A visual inspection should be performed to ensure soil has been removed.

⚠️ **CAUTION:** Do NOT use detergents with ammonia, detergents with abrasives, steel wool, abrasive sponges, steel blades, or cloth with steel thread.

**NOTE:** Use of hydrogen peroxide-based cleaners can discolor the bezel of the digital display. However, this discoloration does not affect the performance, integrity, or usage of the IPV® 1 or display module.

### Stand Assembly

The stand assembly may be cleaned with most mild, non-abrasive solutions commonly used in the hospital environment (e.g., diluted bleach, ammonia, or alcohol solutions). Use a clean, lint-free, non-abrasive cloth for best results. Dry the stand assembly thoroughly after cleaning.
We recommend testing cleaning solutions on a small, non-visible area of the mounting assembly to verify compatibility (will not damage stand).

⚠️ **CAUTION:**
- Do not use strong chemicals or solvents such as acetone or trichloroethylene.
- Do not use steel wool or other abrasive material.
- Never submerge or allow liquids to enter the mounting assembly.
- Wipe away any excess cleaning agent using a water-dampened cloth; do not allow cleaning agent to sit on stand assembly.

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**Breathing Circuit - Phasitron® 5 UC**

It is not necessary to clean the Phasitron® 5 UC breathing circuit after each use. However, rinsing with sterile water is advised. Cleaning instructions are applicable for a single patient using a Phasitron® 5 UC breathing circuit multiple times between applications in same patient.

**NOTE:** When disassembling the Phasitron® 5 UC breathing circuit kit, visually inspect the exterior of all parts, including the tubing, for corrosion, discoloration, pitting, and missing O-rings.

1. Gently rotate nebulizer bowl towards the rear of the Phasitron® 5 until it stops.

2. Separate nebulizer from the Phasitron® 5.

3. While holding the nebulizer cap, twist the nebulizer bowl to remove it from the cap.

Discard any unused liquid in accordance with hospital/institutional protocol.
4. Twist the white cap on the rear of the Phasitron® 5 to remove.

5. Remove the cap.

6. Remove the sliding venturi and spring from the Phasitron® 5 body.

---

**Phasitron® 5 UC Breathing Circuit Cleaning Process**

<table>
<thead>
<tr>
<th>Step</th>
<th>Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Thoroughly rinse each of the disassembled parts (except for tubing harness) under warm running tap water for approximately ten (10) seconds.</td>
</tr>
<tr>
<td>2.</td>
<td>Add fragrance-free liquid soap to a clean bowl or basin filled with warm water.</td>
</tr>
<tr>
<td>3.</td>
<td>Hand wash all parts of the Phasitron® 5 UC breathing circuit kit (including accessories) in the warm soapy water. <strong>NOTE:</strong> Do not wash or submerge the tubing harness.</td>
</tr>
<tr>
<td>4.</td>
<td>Rinse all parts thoroughly using sterile water.</td>
</tr>
<tr>
<td>5.</td>
<td>Gently shake all parts to remove as much water as possible and air dry on a clean, lint-free cloth or paper towel.</td>
</tr>
<tr>
<td>6.</td>
<td>Use a clean, damp cloth to wipe the exterior of the tubing harness using an approved alcohol-based cleaner.</td>
</tr>
<tr>
<td>7.</td>
<td>Reassemble the Phasitron® 5 UC breathing circuit kit and place in a clean bag until the next use.</td>
</tr>
</tbody>
</table>

⚠️ **WARNING:** Do not disinfect the Phasitron® 5 UC breathing circuit kit for reuse for more than one patient.

✨ **NOTE:** The Phasitron® 5 UC breathing circuit is a single-patient device.
11: Troubleshooting

⚠️ CAUTION: If you notice any unexplained changes in the performance of the IPV® 1 driver unit or Phasitron® 5 UC breathing circuit, e.g., if either device is making unusual sounds, or if either device is dropped or damaged in any way, discontinue use and contact authorized service personnel.

<table>
<thead>
<tr>
<th>Problem</th>
<th>Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>No pressure delivery from device</td>
<td>Ensure inlet gas source is on. Check that AIR or O₂ is switched on.</td>
</tr>
<tr>
<td>No percussions</td>
<td>Ensure inlet gas source is on. Check that AIR or O₂ is switched on and AMPLITUDE control is not in the “OFF” position.</td>
</tr>
<tr>
<td>Slow percussion rate</td>
<td>Adjust FREQUENCY knob for higher rate.</td>
</tr>
<tr>
<td>Percussion works, but stalls</td>
<td>Device needs service. Contact authorized service personnel.</td>
</tr>
<tr>
<td>No display</td>
<td>Check batteries are installed and are fully charged. Ensure patient or test lung is attached to the Phasitron® 5 UC. Ensure Amplitude control is not in the “OFF” position and a greater than 7 cmH₂O pressure is achieved.</td>
</tr>
<tr>
<td>Nebulizer not aerosolizing</td>
<td>Ensure a liquid solution is present in nebulizer.</td>
</tr>
</tbody>
</table>

12: Service

IPV® 1 Device

It is recommended to perform an annual preventive maintenance service consisting of a thorough cleaning and a functional evaluation (contact customer service representative).

Stand

Periodically inspect the mounting assembly fasteners. If needed, tighten and adjust the fasteners.

Breathing Circuit - Phasitron® 5 UC

The Phasitron® 5 with Unified Connector breathing circuit is single patient. Clean as instructed in this manual. Replace as necessary.
Service

If the IPV® 1 is not functioning as it should, if it makes unusual noises, or if there are any concerns about device performance or condition, immediately discontinue use. Contact customer service at Customerservice.us@sentec.com.

Changing Display Module Batteries

Displays a low battery indicator when battery capacity is nearing depletion.

1. Press on the display's bezel and twist counterclockwise (left) approximately 20 degrees.
2. Gently pull on the display module's bezel to remove it from the housing.
3. Remove the two old batteries.
4. Install two new CR123A batteries with positive terminals facing the same direction.
5. Wait 30 seconds until the display turns off (blank screen, indicating sleep mode).
6. Install the display back into the housing and twist clockwise until the stop is felt.

NOTE: Do not install display module into the device until the POST test is complete and the screen is blank, indicating sleep mode.

The display module has a USB serial port that is used for factory/service use only.

⚠️ CAUTION: Do not insert any object in the measuring port connection; it may cause irreparable damage.

Disposal of Equipment

At the end of useful life of a unit, dispose of in accordance with local, state, federal, and international laws.

European consumers are obliged by law to dispose of Waste Electrical and Electronic Equipment (WEEE) according to the WEEE Directive 2002/96/EC.

1. All electrical and electronic waste must be stored, collected, treated, recycled, and disposed of separately from other waste.
2. Consumers are obliged by law to return electrical and electronic devices at the end of their service lives to the public collection points set up for this purpose or point of sale. Details to this are defined by the national law of the respective country.
13: Limited Warranty

The manufacturer warrants to the initial purchaser that each new IPV® 1 will be free from defects in workmanship and materials for one year from date of first use (proof of delivery will be required). The manufacturer’s sole obligation under this warranty is to, at its own choice, repair or replace any component – for which the manufacturer acknowledges the warranty coverage – with a replacement component.

Warranty Exclusions and System Performance

Percussionaire® can neither guarantee nor verify product performance characteristics, nor accept warranty or product liability claims, if the recommended procedures are not carried out, if the product has been subject to misuse, neglect or accident, if the product has been damaged by extraneous causes, or if accessories other than those recommended by Percussionaire® are used.

Any serious incident that has occurred in relation to the IPV® 1 system has to be reported to Sentec (regulatory.percussionaire@sentec.com) and/or to the competent authority of the country where the incident occurred. If unsure whether an incident is a reportable event, contact Sentec.
## 14: Technical Specifications

<table>
<thead>
<tr>
<th>Specification</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Dimensions (W x H x D)</strong></td>
<td>23.79 cm x 18.31 cm x 17.53 cm (9.4” x 7.2” x 6.9”)</td>
</tr>
<tr>
<td><strong>Weight</strong></td>
<td>1.45 kg (3.2 lb)</td>
</tr>
<tr>
<td><strong>Operating Range</strong></td>
<td>18°C to 26°C (64.4°F to 78.8°F)</td>
</tr>
<tr>
<td><strong>Storage and Transport</strong></td>
<td>-30°C to +70°C (-22°F to 158°F) Up to 75% relative humidity</td>
</tr>
<tr>
<td><strong>Atmospheric Pressure</strong></td>
<td>Operating: 700 to 1060 hPa Storage: 500 to 1100 hPa</td>
</tr>
<tr>
<td><strong>Gas Source</strong></td>
<td>Wall Gas: 47-80 psi, 3.24-5.5 bar Flow: 35 lpm</td>
</tr>
<tr>
<td><strong>Pulse Interval Ratio</strong></td>
<td>Automatic</td>
</tr>
<tr>
<td><strong>Run Time</strong></td>
<td>Non-continuous</td>
</tr>
<tr>
<td><strong>Aerosol Flow</strong></td>
<td>25 lpm (0.75 - 1 ml per minute liquid consumption rate)</td>
</tr>
<tr>
<td><strong>Pulse Amplitude</strong></td>
<td>0 to 50 cmH₂O/hPa at 100 cycles per minute using Siemens®-style 1L test lung.</td>
</tr>
<tr>
<td><strong>Pulse Frequency</strong></td>
<td>100-300 pulses per minute *Frequency dependent on Gas Supply Pressure and Amplitude setting.</td>
</tr>
<tr>
<td><strong>Mean Airway Pressure (MAP)</strong></td>
<td>0-50 cmH₂O</td>
</tr>
<tr>
<td><strong>Display</strong></td>
<td>Phasitron® 5 with Unified Connector</td>
</tr>
<tr>
<td><strong>Battery Type</strong></td>
<td>Display uses (2) CR123A batteries</td>
</tr>
<tr>
<td><strong>Battery Duration</strong></td>
<td>3,250 Operational hours at 35°C (95°F)</td>
</tr>
<tr>
<td><strong>Accessories</strong></td>
<td>Annual preventive maintenance</td>
</tr>
<tr>
<td><strong>Expected Useful Life</strong></td>
<td>5 years</td>
</tr>
</tbody>
</table>
**Active Mode**: the digital display actively measures the pressure, computes the parameters, and displays them on the display. The Active mode display metrics include pulse frequency rate and mean airway pressure.

**Amplitude**: controls the operating pressure of the entire unit.

**Atelectasis/Atelectatic lung**: a complete or partial collapse of the entire lung or area (lobe) of the lung. It occurs when the tiny air sacs (alveoli) within the lung become deflated or possibly filled with fluid.

**Blended Gas/Air**: the oxygen and air blended according to the prescribed fraction of inspired oxygen (FIO\(_2\)) and delivered to the patient through high-frequency percussive ventilation.

**CMV**: continuous mandatory ventilation.

**Compliant lung**: a measure of the lung expandability, important in ideal respiratory system function, that refers to the ability of the lungs to stretch and expand. Lung compliance can be calculated by dividing volume by pressure.

**Device**: this refers to the IPV® 1 unit.

**Continuous Mandatory Ventilation (CMV)**: the use of a positive pressure ventilator to deliver tidal breaths with conventional target/cycling parameters via an endotracheal tube or tracheostomy tube. Stated in other terms, the tidal volume is generated, in whole or in part, by a mechanical ventilator.

**DISS (Diameter-Index Safety standard)**: a standard for high-pressure gas inlet fittings.

**Entrained air**: the captured ambient air that is automatically drawn in and adjusts to the resistance of the lung.

**Fault Detection**: the digital display has both hardware and software fault detection, a dedicated “watchdog” that runs on an independent clock source and can continue to operate even if the main microprocessor’s clock fails or if the microcontroller pauses in any way.

**Fault Logging**: the digital display software keeps track of several types of hardware and data faults. All faults are logged into the microcontroller’s memory and are retained even if the batteries are removed.

**Fault Mode**: the digital display shows an error message on the display stating “System Failure” and “Contact Factory for Service,” and stays in this mode until both batteries are removed. The displayed information includes the software revision, digital display serial number, total usage time, and an error code.

**Frequency**: the rate of the high-frequency pulses delivered by the IPV® 1.
**FRC (Functional Residual Capacity):** the volume remaining in the lungs after a normal, passive exhalation. The FRC also represents the point of the breathing cycle where the lung tissue elastic recoil and chest wall outward expansion are balanced and equal.

**IPV® therapy:** a type of airway clearance therapy provided by a device that delivers air and aerosol to the lungs at frequencies of 100 to 300 cycles per minute.

**Lung recruitment:** opening up collapsed alveoli to improve oxygenation.

**Mean Airway Pressure (MAP):** the average pressure lungs are exposed to during mechanical ventilation both during inspiration and expiration.

**Nebulizer:** a device for producing a fine spray of liquid, used, for example, for inhaling a prescribed medication.

**Percussive rate:** the high-frequency percussive endobronchial delivery rate of subtidal volumes to the lungs.

**Phasitron® 5:** a unique venturi that acts as a “clutch” mechanism to protect the lung from overpressure. By automatically adjusting to the resistance of the lung, the Phasitron® 5 UC precisely and safely delivers the optimal volume and pressure of air required by the alveolar space.

**Phasitron® 5 UC breathing circuit:** refers to the entire single-patient interface or kit known as the Phasitron® 5 with Unified Connector, including mouthpiece, Phasitron® 5 tubing, and connectors.

**Phasitron® 5 patient-delivery port:** the part of the Phasitron® 5 UC breathing circuit kit that delivers percussive pulses to the patient.

**POST/Power-On Self-Test:** the initial set of diagnostic tests performed by the digital display right after it’s powered on, with the intent to check for any hardware related issues. During this test, the display shows the system information page.

**Proximal Airway Pressure:** pressure measured at the proximal airway (closest to the patient).

**Pulsatile Flow:** the flow created by the Phasitron® 5 is known as Pulsatile Flow, which provides the alveoli with fresh air while gently washing out CO₂ and secretions.

**Report Mode:** mode that displays the session timer and total usage time, alternating with the system information page.

**Usage Timer:** (session timer) the session usage time up to 59 hours and 59 minutes (59:59) before it automatically resets.

**Sleep Mode:** the display is off, but the display module continues to measure pressure at the measuring port. If the display module senses the preset startup pressure, it becomes active.
**System Failure**: when the digital display is in Fault mode, an error message is displayed stating “System Failure” and “Contact Factory for Service.” The display stays in Fault mode until both batteries are removed.

**System Information Page**: when the digital display is in Report mode, the System Information page is displayed for 2 seconds, alternating with the Session Timer. This page displays the copyright, battery life, software revision and the serial number.

**Transient error**: an error that will resolve itself. Most typically these errors manifest as a connection to the database server being dropped. Transient errors can occur, for example, when hardware or network failure happens.

**Usage timer**: a timer that displays the total number of time the device has been operated.

**Wake Mode**: after the POST test, the digital display enters Wake mode and is active for the first 15 seconds of use.