

IPV°-1C USER MANUAL



TRUE-IPV[°] Therapy Device



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The devices and products contained in this manual may be covered by one or more patents.

This manual was originally released and supplied in English. For a list of available translations, contact customerservice@percussionaire.com.

All ventilators should be operated and serviced only by trained professionals. Percussionaire[®] Corporation's sole responsibility with respect to its ventilators, accessories, components, and software, and their use, are as stated in the warranty provided in the manuals. The information set forth herein is believed to be accurate; it is not a substitute for the exercise of professional judgment.

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Chapter 1: Introduction

This chapter provides an overview of the IPV®-1C device and TRUE-IPV® therapy.

Intrapulmonary Percussive Ventilation (IPV®)

Designed specifically for non-continuous institutional/hospital use, the IPV®-1C is a pressure-limited, flow-interrupted and time-cycled ventilator which provides IPV® therapy, a modality of mechanical ventilation, "Intrapulmonary Percussive Ventilation". The IPV®-1C provides high frequency percussive pulses between 100-300 cycles per minute. These high frequency percussive pulses ramify throughout the airways and alveolar ducts augmenting diffusive ventilation in the gas exchange regions of the lungs, allowing improved FRC, CO₂ removal, airway clearance, and lung recruitment.

During operation, the IPV[®]-1C system supplies a continuous dense aerosol mist which is delivered into the lungs during therapeutic percussion, serving to reduce the adhesive and cohesive forces of retained airway secretions.



The IPV[®]-1C, using a Phasitron[®] 5, provides intrapulmonary percussive ventilation either invasively, through an artificial airway, or non-invasively, by mouthpiece, mask, or cannula. The patient may breathe spontaneously, but this is not required.

The IPV®-1C can also be used in-line with conventional mandatory ventilators, delivering TRUE-IPV® therapy and preventing the need to switch between devices.

Operational Pressure Control

Controls the peak operating pressure of the entire unit. This control at maximum output will only provide pressure slightly less than that of the institution. The optimal inlet wall pressure is 50 psi (3.4 bar, 345 kPa).

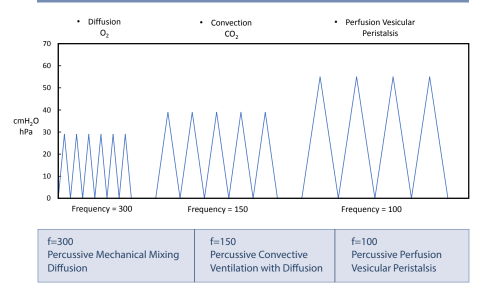
Percussion

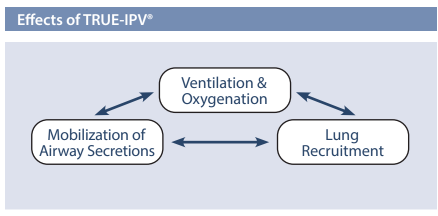
Adjusts frequency of pulses, at a fixed i:e ratio

Manual Inspiration

Delivers a regulated source of gas through the orifice of the Phasitron[®]5 venturi. The longer the button is pressed, the greater the potential for tidal volume delivery.

Three Components of TRUE-IPV®





The effects of TRUE-IPV® therapy occur with or without the cooperation of the patient.

TRUE-IPV[®] provides a percussive gas exchange within the respiratory bronchioles with associated alveolar recruitment maintaining a minimal mean intrathoracic expiratory pressure increase for peripheral lung stabilization. This allows for mechanical ventilation to provide for peripheral lung recruitment while minimizing the potential for induced barotrauma.

Chapter 2: Intended Use

Indications for Use

The IPV[®]-1C is indicated for mobilization of secretions, lung expansion therapy, and the treatment and prevention of pulmonary atelectasis. It can also provide supplemental oxygen when used with compressed oxygen.

Patient Population

IPV®-1C is for use on pediatric and adult patient populations.

Absolute Contraindications

Untreated tension pneumothorax

• Untrained or unskilled operator

Relative Contraindications

History of pneumothorax	Myocardial infarction
Recent pneumonectomy	Vomiting
Pulmonary hemorrhage	• Pulmonary air leak (without functioning chest tube)

Possible Adverse Reactions

Decreased cardiac output	Increased intracranial pressure
Pneumothorax	Increased air trapping
Hyper-oxygenation	Pulmonary air leak
Pulmonary hemorrhage	Hyperventilation
Gastric distension	• Apnea

Physiological Benefits of TRUE-IPV®

Recruitment of atelectatic lung	Mechanical bronchodilation
Improved FRC	May improve breathing pattern
Decreased work of breathing	 Increased secretion mobilization

Clinical Limitations/Restrictions

Use of the IPV®-1C is limited to individuals who have received proper training.

For invasive applications or patients supported by Continuous Mandatory Ventilation (CMV).

WARNING: Due to the therapeutic nature of these devices, they do not have alarms. Consequently, the patient MUST be under continuous observation by a clinician.

WARNING: When used on a patient with an artificial airway (i.e. endotracheal or tracheostomy tube) a clinician must be present so that a one-to-one relationship exists. These devices enhance secretion clearance. Patients must be assessed pre- and post-treatment for a reduced vital capacity/FRC or the need for assistance in clearing airway secretions. Partial deflation of the the cuff during therapy may be necessary; re-inflation per hospital protocol post-therapy.

WARNING: Because pulmonary alveoli cannot be ventilated when their transmitting airways are obstructed, suction should be performed as necessary.

NOTE: A **WARNING** icon indicates a risk of injury to patient or operator. A **CAUTION** icon indicates a risk of equipment damage.

	Type BF Applied Part
	Single Patient Use
Read the manual before use	R Prescription Only
C E marking	REF Catalog Number
Manufacturer	Lot Number
Manufacture Date	European Representative
Non-Sterile	Not Made with
Does Not Contain	Natural Rubber Latex
Plasticizers DEHP, DIBP, DBP, or BBP	Disposal

Document Symbols

Chapter 3: TRUE-IPV® Therapy In-Line with a Ventilator

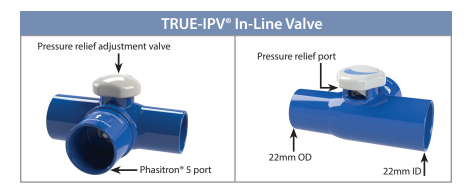


TRUE-IPV[®] In-Line Valve 22mm I.D. x 22mm O.D. Single Patient

Intended Use

The TRUE-IPV[®] In-Line Valve is intended to be used to provide IPV[®] (Intrapulmonary Percussive Ventilation) therapy to intubated patients while assisted by Conventional Mandatory Ventilation (CMV) using pressure-control, volume-control, SIMV-PC etc., when direct connection of IPV[®] is not indicated.

NOTE: For use only with Percussionaire[®] TRUE-IPV[®] ventilator devices. Compatible with all Percussionaire[®] single patient Phasitrons. The IPV[®] In-Line Valve is indicated for patient populations from pediatrics to adults, for whom IPV[®] therapy has been prescribed.



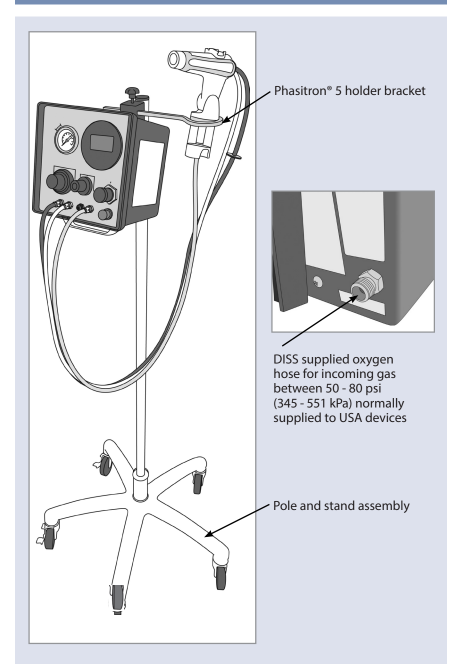
Treatment Frequency

TRUE-IPV[®] in-line use with a ventilator is based on patient need, from 2 times per day up to 6 times per day (every 4 hours), or as recommended by physician. Always use institutional/hospital protocol when possible.

WARNING: Follow institutional protocols before disconnecting ventilator inspiratory limb prior to installation of Percussionaire® TRUE-IPV® In-Line Valve.

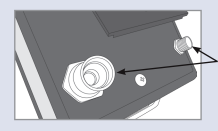
Chapter 4: Setup

Controller and Stand



Rear Panel

Blended Gas/Air Connection

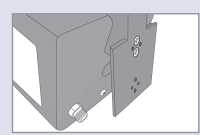


The IPV®-1C can be connected to hospital single gas source or blended gas.

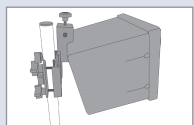
Single or double air/oxygen gas connections available.

Current Gas/Air Connectors Available:			
DISS	USA	NIST	European
AFNOR	French	UNIFOR	Italian
DIN	German	AGA	Scandinavian
BS	British		

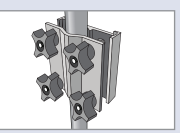
Pole Mount



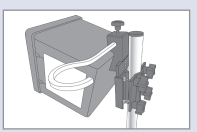
Supplied rear mounting bracket for pole mounting the IPV®-1C device



IPV®-1C device mounted to stand



Adjustable pole mount to attach IPV®-1C device



Side panel holder bracket for convenient storage/placement of the Phasitron[®]5

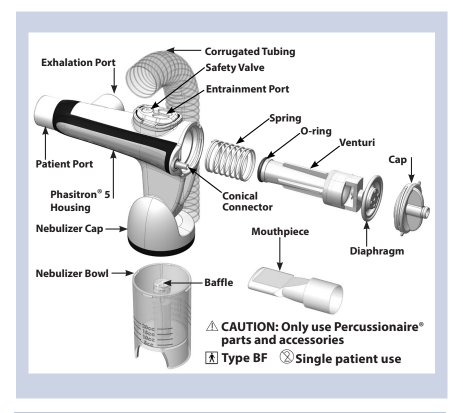
Phasitron[®]5 Breathing Circuit Setup



The patented Phasitron[®] 5 uses a unique venturi as a "clutch" mechanism to protect the lung from over pressure. 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[®]-1C enters the mouth of the venturi. Each air pulse draws up to four times as much additional air into 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.

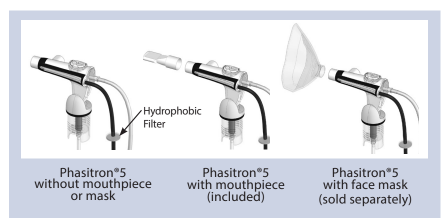
NOTE: TRUE-IPV[®] therapy can only be achieved using the Phasitron[®] 5.

Phasitron[®]5 Diagram

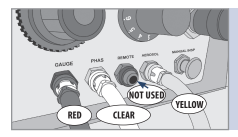


Configurations

Phasitron[®] 5 kit can be used with or without a mouthpiece or standard mask (as shown below). Connection sizes, 15mm ID or 22mm OD.



Connecting to the IPV[®]-1C



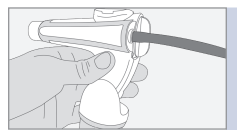
Connect red, clear, and yellow tubing connectors to IPV®-1C controller device.

WARNING: Green remote bulkhead is vented, do not obstruct

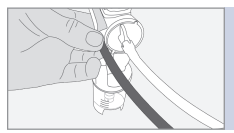
Connect the Tubing Harness to Phasitron[®] 5



Connect yellow tubing quick-connect fitting to nebulizer bowl.

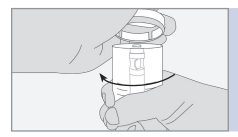


Press red tubing onto conical connector at rear of Phasitron[®] 5 body.

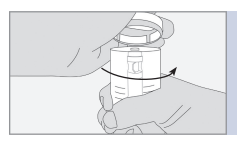


Connect the clear tubing quick-connect fitting to the cap at the rear of the Phasitron[®]5 body.

Adding Saline Solution or Medication



Twist clockwise to open nebulizer bowl. Add saline and/or prescribed medication.



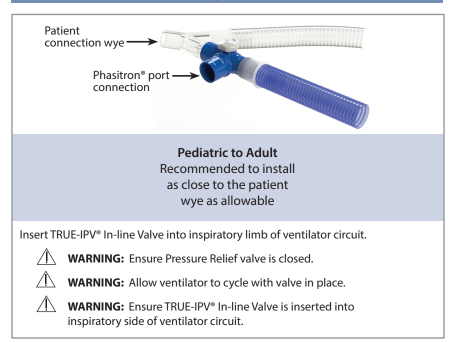
Reverse to close.



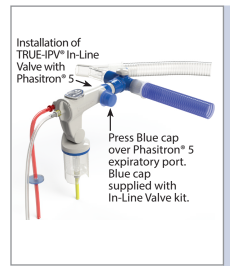
CAUTION: Ensure yellow nebulizer tubing is not bent. This may cause undue stress on connector.

CAUTION: Do not bend nebulizer bowl while holding the tubing. This may cause undue stress on the red line conical connector.

TRUE-IPV® In-Line Valve Setup



Adding Phasitron[®]5 to In-Line Valve



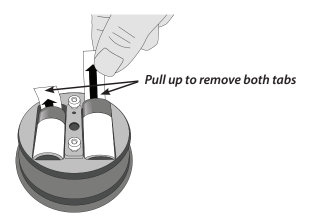
Install provided blue cap (color may vary) onto the Phasitron[®]5 expiratory port. Port must be occluded for proper use and treatment. The cap is provided with TRUE-IPV[®] In-line Valve kit.

Fill nebulizer with 15 to 20 cc normal saline or prescribed medication. Aerosol consumption approximately .75 cc per minute.

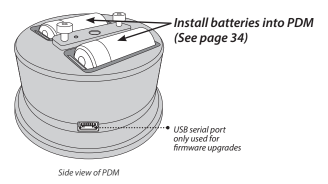
WARNING: Blue cap must be removed when giving a direct treatment, either by mouth, mask or directly connected to endotracheal tube.

Percussionaire® Digital Multimeter (PDM) Setup

NOTE: Remove the PDM from the IPV[®]-1C device, to access battery pull tabs, by turning the PDM counterclockwise.



NOTE: To ensure correct atmospheric pressure calibration at start up, remove batteries, wait 30 seconds and re-install. Allow 15 seconds for power on self test. When screen goes blank, the multimeter can be installed into the device.



NOTE: The PDM has a USB serial port that is used for manufacturing, calibration and firmware upload. It is not enabled during normal operation.

Chapter 5: Controller Functions

Knob, Switch and Button

Knob, Switch, Button	Function
PERCUSSION	The Percussion control knob adjusts frequency of pulses at a fixed i:e ratio. Controls the rate of high frequency volumes delivered. Ranges from 100 up to above 300 cycles per minute. Affects peak and mean airway pressure.
MASTER	The Master Switch turns the controller device ON and OFF. The patient receives TRUE-IPV [®] therapy when the Master Switch is in ON position.
MANUAL INSP PUSH	Manual Inspiration button delivers a regulated source of gas through the orifice of the Phasitron [®] 5 venturi. MarNING: The longer the Manual Inspiration button is pressed, the greater the potential for tidal volume delivery. MarNING: NOT FOR USE WITH NEONATES
OPERATIONAL PRESSURE	The Operational Pressure knob controls the peak operating pressure of the entire unit. The optimal pressure is 40 psi (3.4 bar, 345 kPa) for Pediatrics and Adults.

Percussionaire® Digital Multimeter (PDM)

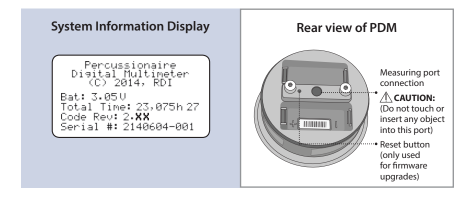


The PDM has six different operating modes: POST, Wake, Active, Report, Sleep and Fault.

Power-On Self-Test (POST) Mode

When batteries are installed in a system, the PDM software displays the software revision, battery voltage, total usage time and serial number for 15 seconds. This Start-Up mode allows the software to perform additional tests on the hardware that are part of the **Power-On Self-Test**. If any errors are detected the PDM enters the Fault mode. It is required that the measurement port be left disconnected and exposed to the atmosphere for the entire duration of the Power-On Self-Test.

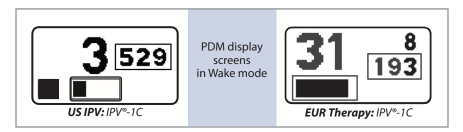
NOTE: Do not install PDM until the POST check is complete and the screen is blank, indicating Sleep mode.



Wake Mode

To wake up the PDM, ensure the ventilator pressure is greater than 2.5 cmH₂O or 2 hPa at the Phasitron^{\circ}5 patient delivery port for more than 1 second.

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



NOTE: *Display numbers are for reference only.*

Active Mode

Model: US IPV Device: IPV[®]-1C Display Metrics: Pulse Frequency Rate, Pulse Amplitude Bar Graph, Mean Airway Pressure, Session Usage Time

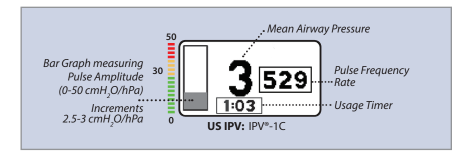
At 16 seconds the PDM enters **Active** mode. The bar graph will change to a numeric display, showing the current usage Session Timer. The display on the right shows the currently measured Pulse Frequency rate.

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

The PDM displays the usage Session Timer in minutes and seconds. The Session Timer is the total time of the current usage. The Session Timer can display a maximum of 59 minutes and 59 seconds. If usage has been stopped for more than 5 minutes, the Session Timer will reset and start over.

The Pulsating Bar Graph on the left side displays Pulse Amplitude calculated as average peak amplitude pressure sample in last 5 seconds, minus amplitude pressure sample in last 5 seconds. The Bar Graph is a visual representation better reflecting AIP and AEP values and represents an estimate of airway pressure. PEEP is represented by a solid bar at the base and AIP is represented by the pulsating peaks of the Bar Graph display.

NOTE: To display most recent usage duration time, see Report Mode.



Model: EUR Therapy **Device:** IPV[®]-1C **Display Metrics:** Pulse Frequency Rate, Mean Airway Pressure, Session Usage Time, Pulse Amplitude Pressure.

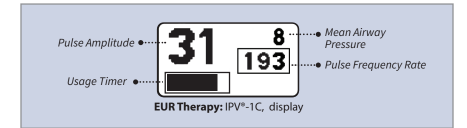
At 16 seconds, the PDM enters Active mode. The bar graph timer will change numeric display, showing the current usage Session Timer. Above the timer reading is the display of the Pulse Amplitude. This is calculated from the pressure measurements at the moment of instantaneous peak and trough amplitude averaged over 5 seconds. The display on the right shows the currently measured Pulse Frequency Rate.

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

The PDM displays the usage Session Timer in minutes and seconds. The Session Timer is the total time of the current usage. The Session Timer can display a maximum of 59 minutes and 59 seconds.

If a usage has been stopped for more than 5 minutes, the Session Timer will reset and start over.

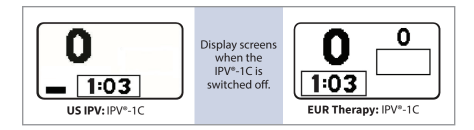
NOTE: To display most recent usage duration time, see Report Mode page 20.



Report Mode

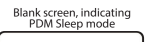
A. 3:5 25h	52	The Session Timer and the Total Usage Timer (A) are displayed for 2 seconds, followed by the System Information page (B) for 2 seconds, alternating. Alternating page display continues for 5 minutes or until usage resumes and the PDM enters Active mode.
B. Percussi Digital Mu (C) 201 Bat: 3.10 v Code Rev: 2.× Serial #: 21	ltimeter 6, RDI	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 flashes (2 seconds on, 2 seconds off) (C) for an additional 25 minutes.
С.		

The PDM enters **Sleep** mode after 25 minutes.



NOTE: When the IPV^{\otimes} -1C is turned off, the measurements will drop to zeros after a few seconds.

Sleep Mode



In **Sleep** mode the LCD 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 2.5 cmH₂O or 2 hPa at the Phasitron[®] patient delivery port, for more than 1 second, the PDM enters the Wake mode.

Fault Mode

System Failure Contact Factory For Service

Code Rev: 2.XX Serial #: 2140604-001 Total Time: 23,075h 27 Err:10/2/3/4/5/6/7/8 The PDM displays an error message on the LCD stating, "Contact Factory for Service" and stays in **Fault** mode until both batteries are removed. The displayed information includes the software revision, PDM serial number, the Total Usage time and an error code for the exclusive use of the factory.

In all other modes, the software continuously monitors the hardware for errors, as well as verifying that each data sample has a valid value. If an error is detected, the software logs the error and reboots the processor, which would cause it 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 a fatal error and the software enters **Fault** mode.

NOTE: Pressure faults are triggered by a continuous pressure of more than 150 cmH_2O for more than 5 seconds during Wake and Active modes.

NOTE: If System Failure screen is displayed, remove batteries for 30 seconds. Replace batteries (note that positive terminals face same direction) and wait 30 seconds until the screen turns off. If POST check runs correctly, PDM may be used. If System Failure screen recurs, contact an authorized Percussionaire[®] service center.

Fault Logging

The software keeps track of several types of hardware and data faults. 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 PDM stops normal operation and enters Fault mode. In this mode, a subset of the collected fault information is displayed on the LCD. This data is intended for manufacturing and repair use only.

The user can exit the Fault mode by removing and replacing the batteries. This resumes normal operation of the PDM but does not erase the faults stored in memory or fix the problem that caused the fault.

Fault Detection

The PDM has both hardware and software fault detection. This is a dedicated hardware "watchdog" that 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.

In addition to the hardware fault detection, the software also implements a fault detection "watchdog". This "watchdog" detects if a software task fails to complete within the specified time, logs an error and resets the processor.

Chapter 6: Pre-Use Check

1.	Connect a Siemens style 1-liter ventilator test lung to the Phasitron [®] 5. (1)
2.	Connect the Phasitron [®] 5 tubing connectors to the IPV [®] -1C using the red, clear and yellow connections. (2)
3.	Rotate the OPERATIONAL PRESSURE knob full counterclockwise P to the stop (OFF) position.
4.	Connect the IPV®-1C to gas source supply.
5.	Rotate the " A " black PERCUSSION knob to the center, up \uparrow position.
6	Rotate the MASTER switch to the "ON" position.
7.	Rotate the OPERATIONAL PRESSURE knob to obtain a pressure between 40-42 psi/2.7-2.9 bar on the operational pressure gauge. Verify pressure by occluding the REMOTE connection for 3 seconds, this will halt percussions and hold the pressure gauge steady. Adjust Operational Pressure knob accordingly.
8.	Rotate the "A" black PERCUSSION knob to the left, full counterclockwise opisition.
9.	Confirm pulse frequency rate of 300 or above.
10.	Rotate the "A" black PERCUSSION knob to the right, full clockwise opposition.
11.	Confirm pulse frequency rate of 100 or below.
12.	Rotate the " A " black PERCUSSION knob to the center, up ↑ position.
13.	Verify that the MAP pressure indicated on the multimeter is greater than 10 cmH $_2$ O.
	NOTE: If using the "EUR" multimeter, MAP is in a different position on display.
	Mean Airway Pressure Mean Airway Pressure Pulse Amplitude Mean Airway Pressure 3 0 3 0 3 0 3 529 1 :03 5 0 1 :03 1 :0

(Continued on p.26)

14.	Occlude green port and observe that pulsations stop.	
15.	. Remove occlusion from green port.	
16.	Press the MANUAL INSP button and observe pressure increase over 40 cmH ₂ O.	
	NOTE: If using the "EUR" multimeter, MAP is in a different position on display.	
17.	Rotate the MASTER switch to the "OFF" position.	
18.	Disconnect gas source supply. Pre-use check is complete.	

Chapter 7: General TRUE-IPV® Therapy Protocol for Adults

	-
1.	Connect IPV [®] -1C to 50-80 psi (3.5-5.5 BAR) gas power source. Master switch is "OFF".
2.	Patient should be in an upright, comfortable armchair or lying with head and shoulders elevated by pillows.
	NOTE: Patient's gravitational position is not a factor with TRUE-IPV [®] .
3.	Auscultate patient for breath sounds, heart and respiratory rate or follow institution guidelines.
4.	Connect Phasitron [®] 5 breathing circuit kit as indicated in <i>Instructions for Use</i> .
5.	Put prescribed medications into nebulizer and add diluent as directed by physician to a maximum of 20 cc. If no medications are prescribed, use normal saline or sterile water as directed by physician.
6.	Rotate FREQUENCY control knob arrow full counterclockwise
7.	Turn IPV®-1C Master switch "ON". Rotate OPERATIONAL PRESSURE control regulator knob for an operating pressure of 30 to 35 psi (2.1-2.4 Bar).
8.	If needed, allow patient to observe and feel pulses from Phasitron [®] 5 on their hand before either connecting to the airway or breathing through the mouthpiece.
9.	When using a mouthpiece, the patient should be instructed to inhale and exhale through the pulses. Most patients will initially allow percussive bursts of air to leak through their nose at the expense of an observable chest movement (wiggle).
10.	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.
	When a patient has an artificial airway, the process is similar. The patient must be observed carefully for signs of distress. While cheek fatigue will be less of a consideration, pauses or breaks may still be necessary for the patient.
11.	Instruct patient to keep their lips and cheeks splinted to avoid nasal air venting. As the patient learns to prevent air from leaking out of the lip seal around the mouthpiece, the PERCUSSION control knob arrow can be gradually rotated clockwise toward the index (12:00).
12.	With some patients, the clinician may have to use an appropriate mechanical airway to administer therapy such as a mask.

(Continued on p.28)

13.	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 briefly rotating the PERCUSSION control knob arrow from EASY to HARD, back and forth (a number of times), then back to the 12:00 arrow position to raise secretions from the bronchial airways.
	NOTE: Adjust treatment to ensure patient comfort.
	As the learning period progresses, the selected source pressure may be increased for effective endobronchial percussion by assessing chest percussion (wiggle).
14.	TRUE-IPV® treatment should continue for 15 to 20 minutes. Recommended daily treatments are 2 to 6 times per day or follow institutional protocol.
15.	When treatment is complete, the IPV®-1C should be turned OFF. The Phasitron® 5 can be rinsed, cleaned and stored in the supplied bag as per hospital infection control policy until the next treatment.
	NOTE: The Phasitron [®] 5 is for a SINGLE patient only, to be used multiple times.
	NOTE: Percussionaire [®] recommends cleaning per your institution's approved practice.

Administering TRUE-IPV[®] Therapy with In-Line Valve

When administering TRUE-IPV[®] Therapy with an In-Line Valve, a pressure control (PC) mode is recommended or follow your institutional protocol.

Mean Airway Pressures (MAP) will increase slightly with the administration of TRUE-IPV[®] In-line therapy with the ventilator. The clinician must be aware of this effect and monitor the patient closely for any adverse side effects.

When using the IPV[®]-1C in pressure-control mode, the in-line valve may remain closed. When using the ventilator in volume-control, the in-line valve may be opened to create a leak.

WARNING: NEVER run device without sterile liquid in nebulizer during treatment. This is required for airway hydration.

A WARNING: Notate the current ventilator alarm and mode settings.

WARNING: Reset occasional CMV high-pressure alarms as they occur. When applying TRUE-IPV[®] in-line, adjust the pressure relief valve to achieve desired Amplitude Pressure per your institutional/hospital protocol. High-pressure alarms should not occur on a regular basis if the Pressure Relief Adjustment valve is set correctly.

NOTE: Patients who are performing T-tube trials or CPAP sprinting may be taken off the ventilator for the IPV® treatment utilizing a flex adapter. Decreasing cuff pressure still applies to this patient population.

NOTE: Following your institutional protocols for cuffed endotracheal tubed patient, the cuff pressure may be lowered.

NOTE: Lowering of the cuff pressure facilitates secretion removal into the oral cavity where they may be suctioned. This also helps in the prevention of tube obstruction in the event copious secretions are mobilized.

NOTE: If chest percussion is inadequate, raise drive pressure (psi gauge) and scan PERCUSSION rate to mobilize secretions.

NOTE: Operational pressure and percussive rate can be adjusted to increase and decrease the amount of chest movement (wiggle).

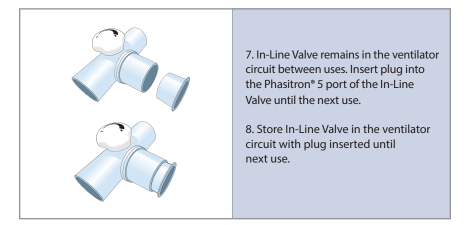
NOTE: Suctioning should be performed as needed.

NOTE: It may take multiple treatments to identify optimal therapeutic effect for each patient.

1.	Ensure the IPV®-1C is "OFF" and connected to a 50 psi/3.2 bar gas source.
2.	Turn incoming air/gas pressure regulator on IPV® device counterclockwise to the stop.
3.	Turn IPV®-1C "ON".
4.	Adjust pressure regulator clockwise, to a starting drive pressure, Pediatric 20 psi/1.4 bar, Adult 30 psi/2.1 bar with a percussive rate of approximately 200 cycles per minute.
5.	Percussion should continue through two complete ventilator cycles to allow ventilator to deliver several machine breaths.
б.	As needed, adjust the pressure relief knob on the In-Line Valve and observe visible chest movement (wiggle).
7.	Monitor patient throughout treatment by observing breath sounds and pulse oximeter for oxygen saturation improvement.
8.	Observe aerosol mist in nebulizer bowl.
9.	Therapy should continue for approximately 15 to 20 minutes, or per institutional/hospital protocol.

Completion of Therapy with In-Line Valve

- 1. If cuff was deflated during treatment, reset cuff pressure.
- 2. Turn off IPV®-1C controller.
- 3. Close Pressure Relief adjustment valve (knob).
- 4. Disconnect Phasitron[®] 5 from TRUE-IPV[®] In-Line Valve and store appropriately.
- 5. Restore ventilator to settings that were present before starting TRUE-IPV[®] treatment.
- 6. Remove cap from Phasitron[®] 5



NOTE: Clean and disinfect In-Line Valve as needed per institutional protocols. In-Line Valve is intended to stay in the ventilator circuit.

Chapter 8: Cleaning and Disinfection

Controller and Stand

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

CAUTION Do not immerse or allow liquids to access the controller.

Clean the controller and stand according to hospital/institutional protocols. Always clean between patients, and when visibly soiled. Clean the controller and stand with a clean, lint-free cloth or paper towel moistened with the cleaner.

CAUTION Use only approved cleaners.

Percussionaire® Digital Multimeter (PDM)

Clean the PDM when visibly soiled or according to facility protocols. Do not spray any type of cleaner directly onto the PDM. Clean the glass with a product or chemical approved for cleaning glass only.

CAUTION Use of cleaning methods not outlined in these instructions may cause damage to the PDM.

WARNING: The cell used in this device may present a risk of fire or chemical burn hazard if mistreated. Do not recharge, disassemble, heat above 100°C (212°F) or incinerate. Replace cell with a CR123A only, or Percussionaire part PRT-B13350. Use of another cell may present a risk of fire or explosion.



Dispose of in accordance with appropriate regulations, country, local and state laws.

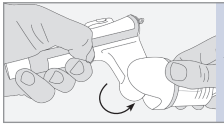
Phasitron[®]5 Breathing Circuit

Follow hospital/institutional guidelines for cleaning and storage between treatments. It is not necessary to clean the Phasitron[®]5 after each use; however rinsing with sterile water is advised. When disassembling the Phasitron[®]5, visually inspect the exterior of all parts, including tubing, for corrosion, discoloration, pitting, and missing O-rings.

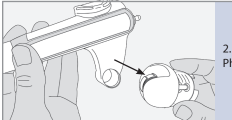
CAUTION Do not immerse the tubing harness.

Disassembly of the Phasitron[®]5

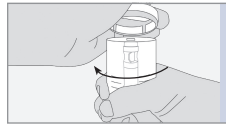
Disconnect tubing from IPV®-1C device and Phasitron®5.



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

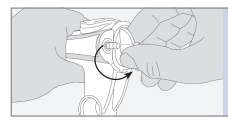


2. Gently separate nebulizer from Phasitron[®]5.

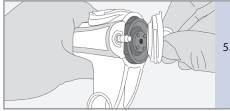


3. Holding the nebulizer cap, twist the nebulizer bowl to remove bowl from cap.

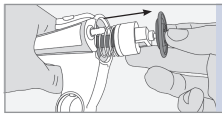
Discard any unused medication in accordance with hospital/institutional protocol.



4. Twist white cap on the rear of the Phasitron[®]5 to remove.



5. Remove cap.



6. Remove sliding venturi with spring from Phasitron®5 body.

Cleaning Phasitron[®]5

1.	Thoroughly rinse each of the disassembled parts (except for tubing harness and filter) under warm running tap water for approximately 10 seconds.
2.	Use fragrance-free liquid soap, added to a clean bowl or basin with warm water.
3.	Hand wash all parts of the Phasitron [®] 5 kit and accessories in the warm soapy water.
4.	Rinse all parts thoroughly using sterile water.
5.	Gently shake all parts to remove as much water as possible, then dry with a clean lint-free cloth or paper towel.
6.	Use a clean, damp cloth to wipe the exterior of the tubing harness with an approved alcohol based cleaner.
7.	Reassemble the Phasitron [®] 5 and place in the supplied bag until next use.
8.	Do not disinfect the Phasitron [®] 5 for reuse with more than one patient.

Cleaning and Disinfecting Solutions

The Phasistron[®] 5 breathing circuit has been tested for biocompatibility with the following cleaning solutions:

Chemical Class	Active Ingredient
Bleach	5.25% Sodium hypochlorite
Alcohol	70% Isopropyl alcohol
Peroxide	3% Hydrogen peroxide
Benzyl Ammonium Chloride	N-alkyl dimethyl ethyl benzyl ammonium chlorides N-alkyl dimethyl benzyl ammonium chloride
Phenolic	Ortho-phenylphenol Ortho-benzyl-para-chlorophenol
Quaternary Ammonium Chloride	Didecyl dimethyl ammonium chloride Alkyl dimethyl benzyl ammonium chloride

Changing PDM Batteries

A Low Battery indicator is displayed when battery capacity is nearing depletion.

1.	Press on the PDM's bezel and twist counterclockwise approximately 20 degrees.
2.	Gently pull on the multimeter to remove it from the housing.
3.	Remove the two old batteries.
4.	Install two new batteries. Note that the positive terminals face the same direction. Wait 30 seconds until screen turns off.
5.	Install the PDM back into the housing and twist clockwise until the stop is felt.
6.	See POST mode instructions to verify display operation.

Chapter 9: Troubleshooting

Problem	Examine	Repair
No pressure indication on the Operational Pressure gauge	Check inlet gas source. Rotate Operational Pressure knob clockwise until 40 psi is indicated.	Connect to Gas Source. Service required.
	Check MASTER switch ON.	Replace or repair MASTER switch.
No percussions	Check inlet gas source.	Connect to Gas Source.
	Check MASTER switch ON/OFF.	Replace or repair MASTER switch.
	Check REMOTE connector for blockage.	Service required.
	Check Operational Pressure regulator.	Service required.
Slow percussion rate	Check REMOTE connector for external blockage.	Rotate percussion knob counterclockwise.
	Percussion knob doesn't change rate.	Service required.
Percussion works, but stalls	Witnessed stall event.	Service required.
No display on PDM	Check battery orientation and strength.	Replace batteries.
	Check both tubing connections.	Reconnect red connections.
	Check if Phasitron® 5 patient port is occluded or connected. to patient.	Occlude patient end of Phasitron [®] 5 if not connected to a patient.
Nebulizer not aerosolizing	Disconnect yellow tubing from IPV®-1C to verify constant flow.	No flow from aerosol connector, service required.
	Check both yellow tubing connections.	Reconnect Yellow Connections.
	Check nebulizer bowl for flow out of nebulizer baffle (Phasitron® 5 diagram).	Clean or replace Phasitron® 5.

CAUTION: If you notice any unexplained changes in the performance of the device, if the device is making unusual sounds, or if the device is dropped or damaged in any way, discontinue use and contact an authorized Percussionaire[®] service center.

Chapter 10: Technical Specifications

IPV®-1C Specifications

Dimensions (W x H x D)	17cm x 24.13cm x 24.13cm (6.7" x 9.5" x 9.5")
Weight	1.99 kg (4.4 lbs)
Operating Range	0°C to 49°C (32°F to 120°F) Humidity < 93% non-condensing
Storage and Transport	-20°C to 60°C (-4°F to 140°F) Humidity < 93% non-condensing
Gas Source	Wall Gas: 50-80 PSI, 3.45-5.5 BAR Flow: 25 LPM
Pulse/Interval Ratio	Automatic
Run Time	Non-continuous
Aerosol Flow	25 LPM
Pulse Amplitude	Digital display, 0 to 99 cmH ₂ O/0 to 97.08 hPa Accurate to $+/-1$ cmH ₂ O 1-5
Pulse Frequency	100-300 pulses per minute
Mean Airway Pressure (MAP)	Digital display
Amplitude Bar Graph	Digital display
Accessories	Phasitron [®] kit P5-10
Battery Type	Multimeter uses (2) CR123A batteries
Required Maintenance	Every 3 years

Phasitron[®]5 Technical Specifications

Size	13.5 mm x 17 mm (5 ¼" x 6 ¾")
Weight	123 g (0.27 lb)
Operating Range	Temp., 0° C to 49° C (32° F to 120° F) Relative humidity range15% to < 90% non-condensing
Storage and Transport	Temp., -40°C to 5° (-40°F to 41°F),
Rate Range	0-999 pulses per minute
Pressure Range	0-150 cmH ₂ O/hPa
Liquid Consumption	.75 cc minute
Safety Valve Release	30-50 cmH ₂ O/hPa
Red Line Filter	1-3 micron hydrophobic
Disposal	Recycle according to local laws
Service Life	6 months or 540 uses whichever is less
Shelf Life	2 years from date of manufacture

Percussionaire[®] Digital Multimeter (PDM) Specifications

Size	73 mm (2.87 inch) diameter
Mass	165 g (0.36 lb)
Storage and	Temp -20° C to 60° C (-4° F to 140° F)
Transport Range	Humidity <93% non-condensing
Operating Range	Temp -20°C to 60°C (-4° F to 140° F), Humidity <93% non-condensing
Display	128 x 64 pixel FSTN chip on glass LCD with reflector
Fault Detection	Independent Hardware and Software Watchdogs
Serial Port	USB (Firmware Upgrade)
Rate Range	50-999 pulses per minute
Pressure Range	1-150 cmH ₂ O/hPa
Pressure Resolution	1 cmH ₂ O/hPa
Pressure Accuracy	Greater of $\pm 0.5\%$ of reading or 1 cmH ₂ O/hPa
Battery Type	CR123A 3.0V (2)
Battery Duration	3,250 Operational hours at 35°C (95°F)
Shelf Life	3.5 Years at 35°C (95°F)

Chapter 11: Service and Repair

Percussionaire[®] recommends an annual Preventive Maintenance (PM) for each device. An annual PM consists of a thorough cleaning, functional evaluation, and if necessary, recalibration. A mandated service is required every 3 years from the date of manufacture or not later than 4 years after first date of purchase. The service includes all new elastomeric components, recalibration, functional evaluation, conformance certification and a one-year warranty on all parts replaced. A device which has not received its required service, whether in use during that period or not, may be considered beyond economic repair. Service by an unauthorized individual or repair maintenance facility will cause the immediate expiration of the clinical readiness of the device. Return the device to Percussionaire[®] or an authorized service center for repair, required service, or annual preventive maintenance.

Disposal of Equipment



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

Chapter 12: Limited Warranty

Percussionaire[®] warrants that the IPV[®]-1C shall be free from defects of workmanship and materials and will perform in accordance with the product specifications for a period of one year from the date of first use (proof of delivery will be required). If the product fails to perform in accordance with the product specifications, Percussionaire[®] will repair or replace – at its option – the defective material or part. Percussionaire[®] will pay customary freight charges to and from Percussionaire[®] or an authorized Percussionaire[®] service center. This warranty does not cover damage caused by non-approved cleaning or sterilization, accident, misuse, abuse, alteration and other defects not related to material or workmanship. Percussionaire[®] disclaims all liability for economic loss, loss of profits, overhead, or consequential damages which may be claimed to arise from any sale or use of this product.

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