INSTRUCTION MANUAL FOR THE NIPPY 3+
POSITIVE PRESSURE VENTILATOR
This book must be kept with the machine

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# NIPPY 3+

## INSTRUCTION MANUAL INDEX

<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Introduction</td>
<td>1</td>
</tr>
<tr>
<td>Description</td>
<td>2-4</td>
</tr>
<tr>
<td>Intended Use</td>
<td>5</td>
</tr>
<tr>
<td>Contraindications</td>
<td>6</td>
</tr>
<tr>
<td>Features</td>
<td>7</td>
</tr>
<tr>
<td>Explanation of Controls</td>
<td></td>
</tr>
<tr>
<td>Fascia Buttons</td>
<td>8-9</td>
</tr>
<tr>
<td>Fascia Display</td>
<td>10</td>
</tr>
<tr>
<td>Outlets</td>
<td>11</td>
</tr>
<tr>
<td>Rear Panel Layout</td>
<td>11</td>
</tr>
<tr>
<td>Explanation of Symbols Used</td>
<td>12</td>
</tr>
<tr>
<td>Getting Started</td>
<td></td>
</tr>
<tr>
<td>The Main Screen</td>
<td>13</td>
</tr>
<tr>
<td>How to Adjust The NIPPY 3 +</td>
<td>13</td>
</tr>
<tr>
<td>The Menu Window</td>
<td>14</td>
</tr>
<tr>
<td>How to use the on-screen Menu</td>
<td>14</td>
</tr>
<tr>
<td>Structure of the Main Menu</td>
<td>15</td>
</tr>
<tr>
<td>Breath Analyser</td>
<td>16</td>
</tr>
<tr>
<td>How to use the on-screen help</td>
<td>17</td>
</tr>
<tr>
<td>Using Help with the Settings Locked</td>
<td>17</td>
</tr>
<tr>
<td>Breathing circuits and masks</td>
<td>18</td>
</tr>
<tr>
<td>Using the NIPPY 3 + Invasively</td>
<td>19</td>
</tr>
<tr>
<td>Using a Humidifier with the NIPPY 3 +</td>
<td>19</td>
</tr>
<tr>
<td>Setting Up</td>
<td></td>
</tr>
<tr>
<td>Setting up the NIPPY 3 + in CPAP Mode</td>
<td>20</td>
</tr>
<tr>
<td>Setting up the NIPPY 3 + in Pressure Support Mode</td>
<td>21</td>
</tr>
<tr>
<td>Setting up the NIPPY 3 + in Pressure Control Mode</td>
<td>22</td>
</tr>
<tr>
<td>Setting up the NIPPY 3 + in IPPV Mode</td>
<td>23</td>
</tr>
<tr>
<td>Setting up the Alarms</td>
<td></td>
</tr>
<tr>
<td>Flow Alarm Auto Set</td>
<td>24</td>
</tr>
<tr>
<td>Flow Alarm Manual Set</td>
<td>24</td>
</tr>
<tr>
<td>Disconnection Alarm</td>
<td>25</td>
</tr>
<tr>
<td>Apnoea Alarm</td>
<td>26</td>
</tr>
<tr>
<td>Max Breath Rate Alarm</td>
<td>26</td>
</tr>
<tr>
<td>Alarm Conditions/Tests</td>
<td></td>
</tr>
<tr>
<td>High Flow</td>
<td>27</td>
</tr>
<tr>
<td>Disconnect</td>
<td>27</td>
</tr>
<tr>
<td>Section</td>
<td>Page</td>
</tr>
<tr>
<td>----------------------------------------------</td>
<td>------</td>
</tr>
<tr>
<td>Low Flow</td>
<td>27</td>
</tr>
<tr>
<td>Mains Fail</td>
<td>27</td>
</tr>
<tr>
<td>High Pressure Fault</td>
<td>27</td>
</tr>
<tr>
<td>Sigh Function</td>
<td>28</td>
</tr>
<tr>
<td>Running on Battery Power</td>
<td></td>
</tr>
<tr>
<td>Battery Run Times</td>
<td>29-30</td>
</tr>
<tr>
<td>Battery Care</td>
<td>31-32</td>
</tr>
<tr>
<td>External battery</td>
<td>33</td>
</tr>
<tr>
<td>Connecting Auxiliary Equipment</td>
<td>34</td>
</tr>
<tr>
<td>Pneumatic Diagram</td>
<td>34</td>
</tr>
<tr>
<td>Specifications</td>
<td>35</td>
</tr>
<tr>
<td>International Standards</td>
<td>35</td>
</tr>
<tr>
<td>Operation Under Extreme Conditions</td>
<td>36</td>
</tr>
<tr>
<td>Accessories and Spares</td>
<td>36</td>
</tr>
<tr>
<td>Warnings and Cautions</td>
<td>37</td>
</tr>
<tr>
<td>Using Supplementary Oxygen</td>
<td>38</td>
</tr>
<tr>
<td>Maintenance</td>
<td></td>
</tr>
<tr>
<td>User Maintenance Schedule</td>
<td>39</td>
</tr>
<tr>
<td>Servicing/Repair</td>
<td>40</td>
</tr>
<tr>
<td>Technical Information</td>
<td>40</td>
</tr>
<tr>
<td>Warranty/Transportation</td>
<td>40</td>
</tr>
<tr>
<td>Factory Service / Repair</td>
<td>40</td>
</tr>
<tr>
<td>Disposal at end of life</td>
<td>40</td>
</tr>
<tr>
<td>EMC Declaration and guidelines</td>
<td>41-43</td>
</tr>
<tr>
<td>Battery Run Time Test Record</td>
<td>44</td>
</tr>
<tr>
<td>Locking the Settings</td>
<td>46</td>
</tr>
</tbody>
</table>

Please note Breas Medical previously traded as B & D Electromedical & some photographs in these instructions still show the name B & D Electromedical.
Introduction

WARNING!

NIPPY 3+ must only be used:

• For the intended treatment in accordance with this operating manual and with the instructions given by the responsible clinical personnel.

• In accordance with the operating conditions specified in this operating manual.

• In original and unmodified shape and only with accessories specified or approved by Breas Medical Ltd.

Every other use may lead to risk of personal injury!

CAUTION!

Read this manual thoroughly so that you completely understand how the NIPPY 3+ is operated and maintained before taking it into use, to ensure correct usage, maximum performance and service life.

WARNING!

This device is not intended for use as a critical care ventilator, transport ventilator or for dependent patients.
Description

The NIPPY 3+ is a pressure controlled, positive pressure ventilator. It compresses ambient air and delivers it to the patient through a full-face mask, nasal mask, nasal pillows or a tracheostomy. The output pressure, timing and alarms can be adjusted by controls on the fascia panel. The Pressure, estimated Tidal Volume, Rate and all settings are displayed on a colour LCD (Liquid Crystal Display) screen.

The screen can be set to dim after a pre-set time (accessed via the user preferences in the main menu). To restore the display, press any button once.

The basic ventilator settings can be achieved using the four buttons to the left of the display. The four buttons to the right of the display adjust the alarms and provide access to more advanced features and adjustments (accessed via a menu).

For greater safety and convenience, the NIPPY 3+ may be equipped with an internal battery. The ventilator is capable of recharging both the internal battery and an external battery when running from the mains electrical supply.

There are 4 modes of ventilation: -

**CPAP**
(Continuing Positive Airway Pressure) Constant positive pressure is applied via the mask. No respiratory support is given in this mode.

**Pressure Support**
IPAP (Inspiratory Positive Airway Pressure) and EPAP (Expiratory Positive Airway Pressure) are set by the physician. The ventilator augments the patient's spontaneous breathing. An adjustable back-up respiratory rate takes over in the absence of an inspiratory trigger from the patient. Ti is limited to a maximum of 3.0 seconds and minimum of 0.7 seconds.

**Pressure Control**
IPAP, EPAP and Ti are set by the physician. A timed inspiration is triggered by the patient’s inspiratory effort. Adjustable back-up rate takes over in the absence of an inspiratory trigger.

**IPPV**
IPAP and Ti are set by the physician. A timed inspiration is triggered by the patient’s inspiratory effort. Patient exhales to atmosphere via an exhale valve fitted in the breathing circuit. EPAP is not used in this mode. Adjustable back-up rate takes over in the absence of an inspiratory trigger.
Alarms

**Power Fail**  If the electrical power to the ventilator is interrupted, an audible alarm will sound. This alarm will run for 5 minutes unless cancelled with the mute button. Once cancelled the power fail alarm will not re-activate.

**Low Internal Battery**  When running on its internal battery, the alarm will operate when the battery is almost depleted.

The user cannot replace this battery. Refer to qualified technical personnel for battery replacement.

**Low External Battery**  When running on an external battery, the alarm will operate when the battery is almost depleted. Machines fitted with an internal battery will automatically switch to internal battery power without alarming.

**Low Pressure**  A pre-set low pressure alarm is provided. If the control pressure falls to below 50% of the set IPAP level for 10 seconds an audible and visual alarm will operate.

**High Pressure**  A pre-set high-pressure alarm is provided. If the pressure rises above 120% of the working pressure, an audible and visual alarm will operate after a 2 second delay.

**Breathing Circuit Disconnect**  A disconnection alarm is provided as a back up to the high flow alarm. If the patient's flow waveform indicates that a significant leak maybe present in the breathing circuit an audible and visual alarm will operate. The sensitivity of the alarm is adjustable.

For an explanation of sensitivity settings see alarm settings, disconnection sensitivity (page 25).

**Breathing Circuit Malfunction**  This alarm warns of a malfunction of the exhale valve in the IPPV mode circuit.

**High Flow alarm**  An adjustable alarm is provided to warn of excess inspiratory flow. This is activated when the inspiratory flow exceeds the set high flow alarm level for 5 seconds. An audible and visual alarm will operate.

**Low Flow alarm**  An adjustable alarm is provided to warn of insufficient inspiratory flow. This is activated when the inspiratory flow fails to achieve the set low flow alarm level for 10 seconds. An audible and visual alarm will operate.

**Apnoea Alarm**  An adjustable Apnoea alarm is provided. If an apnoea is detected an audible alarm will sound with a visual indication on screen.

**Max Breath Rate**  An adjustable maximum breath rate alarm is provided. If the max breath rate is exceeded an audible and visual alarm will operate.

**Fault**  The alarm may also be operated by an internal fault. In this case the fault will be displayed on screen.

*These alarms may be muted for approximately 2 minutes to allow for setting up of the ventilator.*

**Low Internal Alarm/Memory Battery**  An intermittent alarm (short beep) with no onscreen message indicates a depleted mains fail alarm battery. If the ventilator has been stored for more than a few weeks the internal battery will self-discharge. In this case the alarm will stop after the battery has recharged.

The user cannot replace this battery. Refer to qualified technical personnel if the alarm operates when the ventilator is in daily use.
**Estimated Tidal Volume**

The estimated tidal volume is a calculated value, based on time and calibrated flow values. The constant leak through the breathing circuit exhalation port is subtracted from this calculation to give a reasonably accurate estimation of tidal volume. The estimated tidal volume is displayed above the bar graph display.

**Inspiratory Trigger**

The NIPPY 3+ employs flow triggering, detecting the start of the patient’s inspiratory effort when the flow rate exceeds the level set by the Inspiratory Trigger sensitivity.

**Expiratory Trigger**

The expiratory trigger is used in Pressure Support mode only. Towards the end of inspiration, when the inspiratory flow rate drops to the baseline (standing flow caused by exhale port leak) minus the expiratory trigger sensitivity the ventilator will cycle into the expiratory phase.

The inspiratory and expiratory effort required to trigger and cycle the ventilator can be adjusted via the Trigger option in the Menu.

For simplicity, the trigger sensitivity is scaled from 1 – 10, with 1 requiring the least patient effort and 10 requiring more patient effort.
**Intended Use**

The NIPPY 3+ is intended to provide ventilation for non-dependent, spontaneously breathing adult patients with respiratory insufficiency, or respiratory failure.

The device can be used for treating patients non invasively, or for invasive use, with the following interfaces;

- Tracheostomy
- Full Face Mask
- Nasal Mask / Nasal Pillows

The device may be operated in the hospital or home environment.

**IMPORTANT!**

NIPPY 3+ must be prescribed by, and used only under the supervision of a qualified physician.

This manual is only intended for clinical personnel, physicians and trained users who require a working knowledge of the NIPPY 3+.

Alarms must be tested before use and following a change of the breathing circuit as described in the Alarms Conditions/Tests section.

It is the clinicians responsibility to ensure the device is always used with the appropriate breathing circuit, to ensure sufficient CO₂ elimination. All care staff should be trained to ensure that the location of this exhalation port is known and they are aware that this must not be removed, covered or blocked at any time.

Batteries used for power fail back up must be kept in good condition and fully charged at all times. See Battery care section of this manual.
Contraindications

The NIPPY 3+ is not a life support ventilator.

The use of the device may be contraindicated in patients with the following:

- Pneumothorax or pneumomediastinum
- Low blood pressure
- Cerebrospinal fluid leak, recent cranial surgery or trauma, or raised intracranial pressure
- Severe bullous lung disease
- Dehydration
- Risk of vomiting
- An oxygen requirement higher than achievable with 15l/min of entrained low pressure oxygen and the prescribed ventilation settings

Adverse effects

Patients should report any new or unusual chest pain, severe headache or increased breathlessness or increased day time sleepiness to a clinician immediately.

The following side effects may be experienced during the treatment with non invasive ventilation:

- Nasal, mouth or throat dryness
- Bloating
- Ear or sinus discomfort
- Eye irritation from mask leak
Features

1. Optional internal battery operation
2. Comprehensive, auto setting, alarms with mute facility
3. Can be used with Nasal mask, Nasal Pillows, Full face mask or Tracheostomy.
4. User friendly intuitive user interface
5. Easily understood alarm messages displayed onscreen
6. 4 modes of ventilation
7. Universal mains input, operates anywhere in the world without transformers
8. Adjustable flow triggers with trigger indicators
9. Large, colour LCD display, clearly shows all settings
10. 28 days stored, On-screen compliance data
11. Comprehensive event log stores all adjustments, settings, alarm events and user interventions, for download to PC.
12. Breath analyser display, showing pressure and flow waveforms
13. Fast trigger response
14. Minimal maintenance requirement
15. Twelve months parts and labour warranty
16. Auto switching to internal or external battery.
17. Automatic service reminder.
**Explanation of controls**

1. **IPAP** - Selects the Inspiratory Positive Airway Pressure adjustment (scaled in cm H₂O). Value is displayed on screen adjacent to the switch.

2. **EPAP** - Selects the Expiratory Positive Airway Pressure adjustment (scaled in cm H₂O). Value is displayed on screen adjacent to the switch.

3. **Ti** - Selects the inspiratory time adjustment (scaled in Seconds). Value is displayed on screen adjacent to the switch.

4. **Back up** - Selects the Back-up Rate adjustment (scaled in Breaths Per Minute). Value is displayed on screen adjacent to the switch.

5. **Mode** - Displays the mode selection screen

6. -leading - Starts and Stops the ventilator

7.  - Decrements the selected parameter or moves the selection bar down the menu.

8. **Set** - Selects the current menu function displayed by the selection bar. OR double press for battery run time and hours till next service.
9. **+►** - Increments the selected parameter or moves the selection bar up the menu

10. **Mute** - Silences the alarm for 2 minutes. Press and hold for 2 seconds to cancel alarm mute.

11. **Menu** - Displays the menu screen

12. **Help** - Displays context sensitive help messages.

13. **Lo Alarm** - Selects the Low Flow Alarm adjustment (scaled in litres/minute). Value is displayed on screen adjacent to the switch. Changes colour to red in alarm condition.


15. **Ext. Batt** - Indicates that ventilator is running on battery power. This may be internal or external.

16. **Power** - Indicates that external power is connected.

   Ext Batt OFF and Power ON = Battery charging

17. **Start** - Indicates that the ventilator is running.
1. **Mode** - Displays current mode of ventilation.
2. **Pressure Display** - Indicates airway pressure (scaled in cm H₂O). Changes colour to red in under / over pressure alarm condition.
3. **Rate Display** - Indicates patient breath rate (scaled in Breaths Per Minute).
4. **Volume Display** - Indicates estimated inspiratory tidal volume (scaled in millilitres).
5. **Flow Display** - Indicates airway flow (scaled in litres/minute) Changes colour to red in flow alarm condition.
6. **Settings locked symbol** - This symbol shows that the settings are locked.
7. **Alarm Muted symbol** - This symbol shows that the audible alarm has been temporarily silenced.
8. **I Trigger indicator** - “Flashes” each time the inspiratory trigger is initiated by the patient.
9. **E trigger indicator** - “Flashes” each time the expiratory cycle is initiated by the patient.
10. **External battery** - Indicates external battery state of charge, when connected. Red when Discharged.
11. **Internal battery** - Indicates internal battery state of charge. Red when Discharged.
12. **Service Reminder** - major service due
Ventilator Outlets

1. Outlet - Main Air Outlet to breathing circuit
2. EVC Port - Exhale Valve Control outlet
   Connected to the Exhalation valve in IPPV mode
   Leave disconnected in any other mode

Rear Panel Layout

1. Aux. Power - 24 Volt connection for external battery. Connect only recommended batteries, part no 0910
2. RS232 Port - For connection to remote alarm or personal computer. Isolated to 1500 Volts.
3. Power Inlet - Input mains power connector. Double fused and fitted with connector retaining clip.
Explanation of Symbols used on NIPPY 3+ and Accessories

- **Type B Applied parts to EN 60601-1**
- **Alternating Current**
- **Direct Current**
- **Time Delay Fuse**
- **Serial Number**
- **Date of Manufacture**
- **Attention. Consult Accompanying Documents**
- **Switch ON /OFF**
- **Increase Button**
- **Decrease Button**
- **Locked / Unlocked, Purple = total lock, Black = settings locked**
- **Alarm Muted**
- **Battery charged**
- **Battery Discharged**
- **Service Reminder**
- **Dispose of in Line with Local Authority Guidelines**
- **Recycle**
- **Batch code**
Getting Started

To Switch On

Place the NIPPY 3+ on a clean, smooth, hard surface. (NOT carpet)
Connect the power lead to the mains power connector on the rear panel. Plug into
the mains power supply.
Press the Start/Stop button.

To Switch Off

Press the Start/Stop button. The “Switch Ventilator Off” message will appear
onscreen. Press the Start/Stop button again after 2 seconds. There must be a delay
of 2 seconds before each push, to prevent accidental operation.

The Main Screen

The Main Screen is divided into 3 areas
The left-hand side shows the basic ventilator settings, IPAP, EPAP, Ti, BACK-UP,
and Mode, adjacent to its setting button.
The centre section shows the airway pressure, flow, estimated tidal volume and
breath rate.
The right-hand side shows the alarm settings and symbols for alarm mute and locked
settings.

How to adjust the NIPPY 3+

Select the desired parameter with the relevant button.
The reading adjacent to the button will be highlighted by a purple flashing box.
Alter it with the ◀- or ▶+ buttons.
When you have finished, move on to the next adjustment or wait a couple of seconds
for the flashing box to disappear.
E.g. Press IPAP.
IPAP setting will be surrounded by a purple flashing box.
Press +▶ to increase the pressure setting.
The Menu Window

The Main Menu gives access to further adjustments and allows you to view information relating to the ventilator usage.

How to use the on-screen menu

Press the MENU button. The menu window will be displayed in front of the main screen.

Move the selection bar up or down the menu with the ◀- or +► buttons to highlight the desired function and press the SET button.

Follow the on-screen instruction at the bottom of the window

Press MENU at any time to exit and return to the main screen.

Eg.

Press MENU.
Press ◀- button to move the selection bar over “User Preferences”.
Press SET. Press SET again to move the ◀- and +► symbols either side of “Display Contrast”
Press +► to increase contrast - Press MENU to exit.
Structure of the Main Menu

1. Adjust Trigger
   - I Trig View / Adjust
   - E Trig View / Adjust

2. User Preferences
   - User Preferences
     - Alarm Volume View / Adjust
     - Display Brightness View / Adjust
     - Display Contrast View / Adjust
     - Display Dimming View / Adjust
     (Sets time in minutes until display dims, to night time level)
     - Disconnection Alarm View / Adjust
     - Apnoea Alarm View / Adjust
     - BPM Alarm View / Adjust
     - Sigh Settings View / Adjust

3. Compliance Data
   - Total Hours View only
   - Compliance Hours View only
   - Average Daily Use View Only
   Use + or – button to scroll through the data.
   Press and hold + and – buttons to reset compliance data

4. Clear Patient Settings
   Resets machine to default settings and clears the compliance data, ready for a new patient.

5. Warnings & Cautions
   Safety information View only

6. Breath analyser
   Displays Waveforms Enable/Disable
   View Pressure and Flow in graphical form. Press Menu button to return to standard display.

7. Service Menu
   Service Information Restricted Access
Breath Analyser

The breath analyser changes the display to a graphical format, like an oscilloscope. This can be useful to evaluate a patient’s breathing pattern and to assess the effectiveness of the triggering. The pressure and flow waveforms are stored then displayed on the screen.

- The screen displayed shows the stored information from the previous breath.
- The width of the trace is the period of the back-up rate, displayed in seconds on the X axis. E.g. if back up rate is set to 10BPM, the trace width represents 3 seconds.
- The pressure in cm H₂O is displayed on the left Y-axis and relates to the RED trace.
- The flow in litres/minute is displayed on the right Y-axis and relates to the BLUE trace.
- The insp trigger (IT) and exp trigger (ET) points are marked along the X-axis as they occur.
- The display may be “frozen” by pressing the ◄ - button. Press the + ► button to restart the display.
How to use the On-screen Help

Press the HELP button at any time for a list of help topics. Follow the simple on-screen instructions.

Press HELP again to exit.

If during setting up, you require a description of a particular parameter, select it then press HELP. Press HELP again to clear.

Using Help with settings locked

When the settings have been locked, help is limited to a list of more common problems that may arise during use and advice on how to deal with them.

Follow the on-screen instructions
Breathing Circuits

The type of breathing circuit used will depend on the mode of ventilation selected:

Breathing circuits to be assembled in the order A-B-C-D from the ventilator end.

In CPAP, Pressure Support and Pressure Control modes, a single limb circuit with an exhalation port will be required. **DO NOT USE THIS TYPE OF CIRCUIT FOR IPPV MODE**

![Diagram of breathing circuit](image)

**Part Number 0792**
Breathing circuit volume = 570 ml including filter.

In IPPV mode a single limb circuit with an exhalation valve will be required. This type of circuit will have a 22mm diameter main tube and a small tube to connect the exhalation valve to the EVC (exhale valve control) port on the NIPPY 3+. **THIS IS THE ONLY TYPE OF CIRCUIT RECOMMENDED FOR USE IN IPPV MODE**

![Diagram of breathing circuit](image)

**Part Number 0793**
Breathing circuit volume = 600 ml including filter.
**Using the NIPPY 3+ Invasively**

The NIPPY 3+ is safe for use with a tracheostomy or endotracheal tube. An exhalation port must be fitted between the breathing circuit and the tracheostomy fitting. **DO NOT USE THIS TYPE OF CIRCUIT FOR IPPV MODE**

![Diagram of breathing circuit components](image)

**Part Number 0794**

Breathing circuit volume = 620 ml including filter.

**Using a Humidifier with the NIPPY 3+**

Heated breathing circuits are available for use with water bath humidifiers. Follow the instructions supplied with the humidifier.

![Diagram of humidifier usage](image)

**Part Number 0805** for use with a Fischer and Paykel Humidifier

Breathing circuit volume = 1290 ml including filter and water bath.
Setting up the NIPPY 3+ in CPAP mode

1. Place the NIPPY 3+ on a clean, level surface.
2. Connect the breathing circuit tube to the outlet. It is recommended that a bacterial filter be fitted between the outlet and the 22mm diameter breathing tube.
3. The breathing circuit must contain an intentional leak to prevent re-breathing of exhaled air. This may be achieved with either an integrated leakage in the mask or a leakage/exhalation port.
4. Connect the mask to the outlet tubing on the breathing circuit.
   **Note:** Do not fit the mask onto the patient at this point.
5. Press the Start/Stop button and select CPAP mode.
6. Set the High Alarm to 160 lpm and the Low Alarm to 20lpm. This will help to stop the alarm becoming a nuisance during setting up. Alarm settings will be finalised later.
7. Set EPAP to the prescribed pressure.
8. The patient may now **hold** the mask to the face.
9. Allow the patient to get used to the interface then-fasten the interface to the patient.
10. Set Up the alarms, as detailed in the “Setting Up the Alarms” section of this manual
    - **Test the alarms as detailed in “Alarm Conditions and Tests” section of this manual.**
11. Lock the settings to prevent unauthorised adjustment. See back page.

Remove the patient interface or disconnect the patient circuit from the outlet port, before switching off.
Setting up the NIPPY 3+
in Pressure Support mode

1. Place the NIPPY 3+ on a clean, level surface.
2. Connect a clean or new breathing circuit tube to the outlet. It is recommended that a bacterial filter be fitted between the outlet and the 22mm diameter breathing tube.
3. The breathing circuit must contain an intentional leak to prevent re-breathing of exhaled air. This may be achieved with either an integrated leakage in the mask or a leakage/exhalation port
4. Connect the mask to the outlet tubing on the breathing circuit.
   Note: Do not fit the interface onto the patient at this point.
5. Press the Start/Stop button and select Pressure Support mode.
6. Set the High Alarm to 160 lpm and the Low Alarm to 20 lpm. This will help to stop the alarm becoming a nuisance during setting up. Alarm settings will be finalised later
7. Ensure the settings are adjusted as prescribed.
8. The patient or physician if required may now hold the interface to the face.
9. Allow the patient to get used to the interface. Objective assessments should verify the settings are suitable and clinically effective.
10. If the inspiratory or expiratory trigger needs to be adjusted, select “Adjust trigger level” from the menu and adjust to suit the patient and to ensure optimal patient synchrony.
11. Read the breath rate from the display (top of screen). This should match the value observed when assessing the patient. If the rate has increased, make sure that the trigger is not so sensitive that it is causing “self triggering”.
12. Allow the patient to get used to the interface then-fasten the interface to the patient.
13. Set Up the alarms, as detailed in the “Setting Up the Alarms” section of this manual
   - Test the alarms as detailed in “Alarm Conditions and Tests” section of this manual.
14. Check the rate and alarms when the patient is asleep. The inspiratory and expiratory trigger indicators should “flash” at the beginning of each breath.
15. Lock the settings to prevent unauthorised adjustment. See back page.

Remove the patient interface or disconnect the patient circuit from the outlet port, before switching off.
Setting up the NIPPY 3+ in Pressure Control mode

1. Place the NIPPY 3+ on a clean and level surface.
2. Connect a clean or new breathing circuit tube to the outlet. It is recommended that a Bacterial filter be fitted between the outlet and the 22mm diameter breathing tube.
3. The breathing circuit must contain an intentional leak to prevent re-breathing of exhaled air. This may be achieved with either an integrated leakage in the mask or a leakage/exhalation port.
4. Connect the interface to the outlet tubing on the breathing circuit.
   **Note: Do not fit the interface onto the patient at this point.**
5. Press the Start/Stop button and select Pressure Control mode.
6. Set the High Alarm to 160 lpm and the Low Alarm to 20 lpm. This will help to stop the alarm becoming a nuisance during setting up. Alarm settings will be finalised later.
7. Ensure that the settings are adjusted as prescribed.
8. Allow the patient to get used to the interface. Objective assessments should be performed to establish effective ventilation is achieved.
9. Adjust the Inspiratory time (Ti) to ensure adequate ventilation and patient comfort.
10. If the Inspiratory trigger needs to be adjusted, select “Adjust trigger level” from the menu and adjust to achieve optimal patient synchrony.
11. Read the breath rate from the display (top of screen). This should match the value observed when assessing the patient. If the rate has increased, make sure that the trigger is not so sensitive that it is causing “self triggering”.
12. Allow the patient to get used to the interface then-fasten the interface to the patient.
13. Set Up the alarms, as detailed in the “Setting Up the Alarms” section of this manual.
   - Test the alarms as detailed in “Alarm Conditions and Tests” section of this manual.
14. Check the rate and alarms when the patient is asleep. The inspiratory trigger indicator should “flash” at the beginning of each breath.
15. Lock the settings to prevent unauthorised adjustment. See back page.

Remove the patient interface or disconnect the patient circuit from the outlet port, before switching off.
Setting up the NIPPY 3+
in IPPV mode

1. Place the NIPPY 3+ on a clean, level surface.
2. Connect a clean or new breathing circuit tube to the outlet. It is recommended that a Bacterial filter be fitted between the outlet and the 22mm diameter breathing tube.
3. Connect the small exhalation valve tube to the EVC outlet.
4. Connect the mask to the outlet tubing on the breathing circuit.

Note: Do not fit the interface onto the patient at this point.

5. Press the Start/Stop button and select IPPV mode.
6. Set the High Alarm to 160 lpm and the Low Alarm to 20 lpm. This will help to stop the alarm becoming a nuisance during setting up. Alarm settings will be finalised later.
7. Ensure that the settings are adjusted as prescribed.
8. The patient or physician may now hold the interface to the face.
9. Allow the patient to get used to the interface. Objective assessments should be carried out to verify optimal ventilation is being achieved.
10. Adjust the Inspiratory time (Ti) to ensure adequate ventilation.
11. If the Inspiratory trigger needs to be adjusted, select “Adjust trigger level” from the menu and adjust to ensure optimal patient synchrony.
12. Read the rate from the display (top of screen). This should match the value observed when assessing the patient. If the rate has increased, make sure that the trigger is not so sensitive that it is causing “self-triggering”.
13. Set Up the alarms, as detailed in the “Setting Up the Alarms” section of this manual.

Test the alarms as detailed in “Alarm Conditions and Tests” section of this manual.

14. Check the rate and alarms when the patient is asleep. The inspiratory trigger indicator should “flash” at the beginning of each breath.
15. Lock the settings to prevent unauthorised adjustment. See back page.

Remove the patient interface or disconnect the patient circuit from the outlet port, before switching off.
Setting Up the Alarms

Flow Alarm Automatic Setting

For convenience, the high and low flow alarms may be set automatically. Varying leak or the patient getting out of synch with the ventilator may disrupt the measurement process. Therefore, it is vital that the alarms are tested after setting up.

Low Flow Alarm
- Press and hold the Low Flow Alarm button
- The ventilator will read the peak flow, the inspiratory baseline or leak flow and set the low flow alarm to the mid-point between these two values.
- This setting may be further adjusted manually if required.
- Test the alarm as detailed in “Alarm Conditions and Tests” section of this manual.

High Flow Alarm
- Press and hold the High Flow Alarm button
- The ventilator will read the peak flow and set the high flow alarm to this value plus 30% or 20 l/min, whichever is greater.
- This setting may be further adjusted manually if required.
- Test the alarm as detailed in “Alarm Conditions and Tests” section of this manual.

Warning!
The auto set facility is not infallible. Varying leak can cause erroneous readings. The settings must be verified and the alarm function tested. If the alarm settings are not correct or the alarms do not operate when tested, proceed to the manual set up.

Flow Alarm Manual Setting

Low Flow Alarm
- Note the peak flow reading. This is the patient’s peak inspiratory flow.
- Disconnect the breathing circuit at the mask or tracheostomy and occlude the end (do not obstruct the exhale port). Note the peak flow reading. This is the leak flow.
- Reconnect the breathing circuit.
- Set the Lo alarm to a value approximately half way between the leak flow and the peak inspiratory flow.
- Test the alarm as detailed in “Alarm Conditions and Tests” section of this manual.
- If the patient is being treated via a nasal mask, this alarm is not required and may be set to minimum.

High Flow Alarm
- Note the peak flow reading. This is the patient’s peak inspiratory flow.
- Disconnect the breathing circuit at the mask or tracheostomy end. Note the peak flow reading. This is the disconnected flow.
- Set the Hi alarm to a value a little higher than the peak inspiratory flow. This must be lower than the disconnected flow. Allow for an increase in flow if the patient takes a deep breath.
- Test the alarm as detailed in “Alarm Conditions and Tests” section of this manual.
**Disconnection Alarm**

The disconnection alarm acts as a back up to the high flow alarm.

The inspiratory and expiratory flow waveform of the patient is analysed. If the waveform indicates that a significant leak maybe present in the breathing circuit an audible and visual alarm will operate after 10 seconds.

There are three different sensitivity levels or it may be disabled if not required. Select the appropriate sensitivity level for your patient. Press SET to store the setting.

It is essential that the high flow and disconnection alarms are tested with the circuit and interface/tube used by the patient to ensure the device alarms appropriately - see page 27 testing alarms.

![Disconnection Alarm](image)

**High Sensitivity** – The alarm will be activated during the detection of a leak which is less than that required to trigger the standard sensitivity setting. This may be useful for very restrictive breathing circuits or small tracheostomy tubes.

**Standard Sensitivity** – Default level.

**Low Sensitivity** – The alarm will be activated during the detection of a leak which is greater than that required to trigger the standard sensitivity setting. This may be useful where mask leaks cause nuisance disconnect alarms. Verify that patient is properly ventilated during these periods before using the Low setting.

**Disabled** – Disconnect alarm is switched off. Useful where mouth breathing or excessive leak causes nuisance disconnect alarms. Verify that patient is properly ventilated during these periods before using this setting. The High Flow alarm will activate if a flow exceeds the set threshold. A full risk assessment must be carried out and documented before activating this setting, and must only be set by the prescribing physician.
Apnoea alarm

Select USER PREFERENCES from the main menu. From the OPTIONS MENU, select APNOEA ALARM.

An apnoea event is defined as a cessation of spontaneous breathing for a set time delay. The apnoea alarm will be activated if this condition is detected. Ventilation will continue at the backup rate during apnoea events.

To activate the apnoea alarm, select ON and select the required time delay. The total number of apnoea events during the last seven days or since reset, will be displayed along with the time duration of the longest single apnoea. Press + and – together to reset the apnoea count.

Max Breath Rate Alarm

The max rate alarm will be activated if the patients breath rate exceeds the set level. Select USER PREFERENCES from the main menu. From the OPTIONS MENU, select RATE ALARM. To activate the rate alarm, select ON and set the required max rate.
Alarm Conditions/Tests

Test the alarms prior to use or daily for machines that are in constant use. Before testing alarms, ensure that the alarm is not muted. To cancel the Mute, press and hold the mute switch until a beep is heard (2 seconds).

**High Flow Alarm**

If the inspiratory flow exceeds the high alarm value, the alarm will be activated accompanied by the on-screen, high flow alarm message.

**To Test** Disconnect the breathing circuit at the patient interface and allow the flow to exceed the set alarm value. The alarm will be activated after a delay of 5 seconds, accompanied by the on-screen, high flow alarm message.

**Disconnect Alarm**

The disconnect alarm acts as a back-up to the high flow alarm to warn of breathing circuit disconnection.

**To Test** Switch on the ventilator and cancel the Mute. Disconnect the breathing circuit at the patient interface. The High Flow alarm will be activated after a delay of 5 seconds and will be replaced by the disconnect alarm after 10 seconds, accompanied by the on-screen disconnect alarm message.

**Low Flow Alarm**

The low flow alarm warns of insufficient inspiratory flow. This could be caused by a blockage in the patients’ airway or breathing circuit. The resulting drop in flow will activate the alarm.

**To Test** Switch on the ventilator and cancel the Mute. Occlude the outlet and wait 10 seconds. The alarm will sound, accompanied by the low flow alarm message and the flow display will turn red.

**Power Fail Alarm**

If both mains and internal power to the NIPPY 3+ fails, the alarm will operate and continue for approx. 5 minutes. Press the mute button to silence the alarm.

**To Test** Machines NOT fitted with an internal battery - start the NIPPY 3+ and switch off the mains power at the wall socket. The screen will go blank after a few seconds and the alarm will sound. Press the mute switch or restore the power and re-start the ventilator to silence.

Machines fitted with internal battery – Power fail alarm will only operate if the internal battery is completely discharged and the mains fails. The only way to test it is to run the ventilator on battery power until the battery is exhausted and the power fails.

**High Pressure Alarm**

If the airway pressure exceeds 120% of the IPAP setting for more than 2 seconds, the alarm will sound and the pressure display will turn red.

*It is not possible for the user to test this function.*

**Fault Alarm**

The fault alarm indicates a fault in the machine. The on-screen message will indicate the nature of the fault. The message may be temporarily hidden by pressing the Mute button. It is not possible for the user to test this function.

*If you receive a fault message at any time, DO NOT continue to use the ventilator. The machine MUST be referred to suitably qualified technical personnel for investigation/repair.*
Sigh Function

The ventilator can be programmed to deliver an occasional sigh with a larger tidal volume. This prevents collapse of the alveoli (atelectasis) which can result from the patient constantly inspiring the same volume of gas. Sigh is available in all modes except CPAP.

The Sigh function provides a timed inspiration at the IPAP and TI settings specified in the Sigh Settings screen.

Sigh may be set for cyclic operation, sigh breath at the specified interval or an external switch (Manual mode). Or a combination of both, where the cyclic sigh may be supplemented by extra sigh breaths on demand.

Setting the Sigh function

Press MENU and select USER PREFERENCES. In the User preferences screen select SIGH SETTINGS.

Use the SET button to scroll down the settings, using the +/- buttons to alter the settings.

Sigh Interval sets frequency of cyclic sigh
Ext Trig = manual switched sigh

Symbol ■ shows Manual Sigh function is enabled.
Symbol □ shows Cyclic Sigh function is enabled.
Blue Symbol = Sigh mode enabled.
Green Symbol = External sigh trigger acknowledged.
Purple Symbol = Sigh breath activated.
Sigh breath pressure (S) is indicated on the pressure bar-graph display.

Note: If the ventilator IPAP is adjusted after the sigh has been set up, this may affect the sigh setting.

If the IPAP is reduced the sigh will be reduced by the same amount, to maintain the difference between IPAP and Sigh. This can be overridden in the Sigh Settings screen.

If IPAP is increased to a value equal to the sigh setting, the sigh value will “track” the IPAP. This effectively makes IPAP and Sigh the same. This is done to prevent accidental increase of the sigh breath. The sigh will need to be reset in the Sigh Settings screen. Sigh is cancelled when Mode is changed.

The IPAP during Sigh cannot be lower than normal IPAP.

Regardless of the back-up rate, a sigh breath is timed with an IE ratio of 1:2. This does not affect triggering.

High flow and disconnect alarm is suspended during Sigh.

No more than 5 consecutive sigh breaths are allowed.
Running the NIPPY+ on Battery Power

The NIPPY + may be powered from the mains, internal or external battery.

IMPORTANT BATTERY CARE RULES
1. Regularly check battery run times for your system at your prescription settings and record in the log.
2. If you have more than one ventilator, check this one too.
3. Never check the on-screen run time estimate when running on mains power.
4. Failure to perform regular battery performance checks could result in failure during loss of mains power. Check Monthly.
5. Know the difference between battery run time and battery state of health. The run time is the running time expected from a healthy battery. The state of health is the condition of the battery in terms of its total service life.

Power Source Priority
1. - Mains Electricity
2. - External Battery (if connected)
3. - Internal Battery (if fitted)

In order to save battery power, the ventilator will always run on mains electricity if it is present.

If the mains fails or is not connected, the ventilator will select the external battery as the next choice. The external battery, if present, will always be run flat before the internal battery is selected.

If there is no external battery, the ventilator will switch to its internal battery.

If there is no internal battery the ventilator will shut down and alarm to warn of power failure.

Battery Run Times

Battery run times are dependent on the battery age, condition, ambient temperature, ventilator settings and amount of leak. High pressures and /or high breath rates use more power and therefore shorten run times. Large leaks use more power and shorten run times.

Run times will gradually reduce as the battery ages regardless of how much use the battery has had.

Battery run times will typically be 20% more in IPPV mode because of the absence of the exhale port leak.
Battery Run Time Estimate Screen

Double press the SET button to display a bar graph of run time. The time for each battery connected will be calculated, according to the ventilator settings, and shown on the screen.

This is an indication of the battery state of charge not its health. For most accurate result, run the ventilator on battery power for 5 minutes prior to this check.

The results shown assume that the connected battery is in good serviceable condition. It is not possible to take into account the state of health of the battery. The only way to determine the state of health of the battery is to perform a run time test.

When checking expected battery run time the ventilator must NOT be connected to the mains.

Low Battery Alarm

Low battery = alarm sound, and on screen message, “Battery Power Running Low” and flashing red battery icon.

Flat Battery = alarm sound and on-screen message “Battery Power Exhausted. About to switch off. Connect to Mains Power Now” and flashing red battery icon.

Critical battery = 10 minutes after battery exhausted warning, alarm sound continuous tone and auto power off.

After auto power off, the constant alarm will continue until the Mute is pressed.

Charging the batteries

Internal and external batteries are automatically charged when the ventilator is connected to the mains supply. Charging takes approximately 8 - 11 hours depending on ventilator settings.

Charging priority is given to the internal battery. Charging of the external battery if connected will begin after the internal battery is charged. When charging is complete the batteries will be monitored to maintain charge.
Battery Care

- Check the running time of your system regularly.
- Always make sure batteries are fully charged before use.
- The battery should be recharged as soon as possible after use.
- This type of battery does not suffer from the memory effect that is widely talked about and does not need to be fully discharged before charging.
- Batteries like to be used. A new battery may require several charge/discharge cycles before it reaches its maximum performance. The same applies to a battery that is only used occasionally with long periods in storage.
- If you have more than one external battery, use them in rotation.
- If you are in doubt regarding the state of charge, charge for at least 24 hours.

Battery Life
The battery will deteriorate due to age and usage. It is not possible to predict the ageing. The life span of the battery will depend on a number of factors, including:
- Age, number of charge cycles, depth of discharge, charge and discharge temperature.
- An internal battery that is not used will age prematurely.
- An internal battery that is heavily used will age gradually and lose its capacity.
- An internal battery that is moderately used will last longest. It has been designed to be used.
- When a battery is towards the end of its life, the run times will be much shorter. This is why it is important to carry out a run time check regularly.
- The end of life is defined by the maximum running time falling to 75% of that of a new battery (refer to the Battery Testing section.)
- Service life is 2 years under normal operating conditions. Replace the battery when running times drop below those indicated or after 2 years.
- If the battery icon remains RED after several hours charging there may be a fault with the battery of the ventilator.

Battery Storage
- This type of battery is best stored partly charged.
- A battery that is not in use will slowly discharge. This rate of discharge increases with temperature.
- Optimal storage temperature is between -20°C and +20°C. It must be below 40°C.
- After storage in a cold environment allow 24 hours for the battery to reach room temperature before use.
- Fully charge the battery before use.

Battery Run Time Test (Health test)
- Ensure the battery is fully charged.
• Run the ventilator from the battery until the low battery alarm operates and record the running time. Look up the run time in the table. If the battery is not achieving minimum run time replace it.
• If the battery is good, fully recharge it immediately after testing.
• A run time record is provided at the end of this manual.

Table of typical running times
Pressure Control / Support mode with EPAP set to minimum @ 20BPM

<table>
<thead>
<tr>
<th>IPAP</th>
<th>Run Time (Hours)</th>
<th>Min Run Time (Hours)</th>
</tr>
</thead>
<tbody>
<tr>
<td>15</td>
<td>11</td>
<td>8.25</td>
</tr>
<tr>
<td>20</td>
<td>9.5</td>
<td>7.25</td>
</tr>
<tr>
<td>25</td>
<td>8.5</td>
<td>6.25</td>
</tr>
<tr>
<td>30</td>
<td>6.5</td>
<td>5</td>
</tr>
<tr>
<td>35</td>
<td>5.5</td>
<td>4</td>
</tr>
<tr>
<td>38</td>
<td>5</td>
<td>3.75</td>
</tr>
</tbody>
</table>

Battery Health Quick Test

If it is not practical to check battery run times, a crude quick test may be performed to identify a battery that is at the end of its life and has not been noticed by the user or for a quick check on a battery that has not been used regularly. This test is not an indication that the battery is good, but is a useful way of identifying a failed battery. This test is not intended to replace regular run time testing used to determine battery state of health.

Quick Test Procedure
1. Remove the external battery from the Aux Power socket if being used.
2. Connect NIPPY+ to mains power supply and fully charge the internal battery overnight.
3. Turn the NIPPY+ on with the patient circuit fitted to the outlet.
4. Remove the patient circuit from the outlet and allow the NIPPY+ to ‘free flow’. This will create maximum flow and load on the battery.
5. If the battery symbol turns red or the NIPPY+ shuts down within 2 minutes the battery MUST be replaced immediately.

Disposal of depleted batteries
Depleted batteries may be disposed of in line with local authority regulations.
External Battery

An external battery may be used to increase the running time. This battery will power the NIPPY+ for the same time as an internal battery, depending on settings and leak.

External Battery, part number 0910.
External battery charger, part number 0911
These batteries should never be used to run any other type of equipment.

Instructions for Use

• Connect the battery to the NIPPY+ Aux Power input.
• Switch on the NIPPY+. The Ext Batt light will flash and a “Running on battery power” message will be displayed on the NIPPY+ screen. Press the alarm mute button to hide the message.
• To disconnect a battery: Always switch off the ventilator first. Press the plug release button on the connector and withdraw the connector.

To Charge an External Battery

• Connect the battery to the NIPPY+ whilst it is connected to mains electricity.
Alternatively, charge with the battery charger as follows:-
• Place the charger on a smooth flat surface.
• Connect the charger to the battery socket before switching on the mains power.
• Connect the mains plug to the AC supply and switch on.
• Leave on charge until the charged / ready indicator lights.
• Batteries may be left connected to the charger until required for use.

Safety

• Warning! High voltages exist inside the charger.
• Do not remove the cover. Return to Breas Medical Ltd if a fault occurs.
• Do not expose to water or dust.
• Do not cover the charger whilst in use
• Do not use any other type of battery charger. This could lead to damage to the battery and personal injury.
• Do not attempt to connect any battery other than those supplied by the manufacturer. Use of any other type of battery could lead to personal injury and damage to the ventilator.
• Ensure that the mains lead is not damaged.
• Batteries may produce explosive gases during charging. Always charge away from sparks or sources of ignition. Do not smoke near a battery whilst charging.
• Disconnect from the mains before disconnecting the battery from the charger
Connecting auxiliary monitoring equipment

For monitoring or downloading data the NIPPY 3+ may be connected to a PC or Laptop computer. The NIPPY 3+ isolated RS232 port is safe for use with any domestic PC or laptop computer. However, when assembling a system, the completed system should comply with EN60601-1 (medical systems). For example, most computers do not comply with this standard, so it should be sited at a distance, which makes it impossible to touch the computer and the patient at the same time.

The RS232 port may also be used to run a remote alarm unit.

Ventilator System Pneumatic Diagram

1. Fresh air inlet
2. Blower (compressor)
3. Flow Sensor
4. Outlet connector
5. EVC Port (IPPV mode only)
6. Breathing Circuit
7. Exhalation Port
8. Patient Connection Port
9. Exhalation Valve (IPPV mode only)
## SPECIFICATIONS

<table>
<thead>
<tr>
<th>Specification</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Supply Voltage</td>
<td>100 - 240 V alternating current</td>
</tr>
<tr>
<td>Supply Frequency</td>
<td>47 - 63 Hz</td>
</tr>
<tr>
<td>Maximum Input Current</td>
<td>0.40 – 1.0 Amperes</td>
</tr>
<tr>
<td>Fuse Ratings</td>
<td>2 x T 1.6 A 20mm</td>
</tr>
<tr>
<td>Dimensions (mm)</td>
<td>Length - 297 Width - 223 Height - 132</td>
</tr>
<tr>
<td>Weight</td>
<td>4.5 kg</td>
</tr>
<tr>
<td>Ambient Operating Temperature</td>
<td>40°C, 104°F Max</td>
</tr>
<tr>
<td>Digital Output</td>
<td>RS232 Isolated to 1500 Volts</td>
</tr>
<tr>
<td>All displayed readings expressed as</td>
<td>ATPD</td>
</tr>
<tr>
<td>Max. Output Pressure</td>
<td>38cm H₂O (44cm fault condition)</td>
</tr>
<tr>
<td>Calibrated pressure Range</td>
<td>0 - 38cm H₂O</td>
</tr>
<tr>
<td>Accuracy of pressure reading</td>
<td>+/- 3.0% F.S. +/-1% zero</td>
</tr>
<tr>
<td>Max. Output Flow</td>
<td>200 L/min. (unrestricted)</td>
</tr>
<tr>
<td>Max Volume Reading</td>
<td>2000 millilitres</td>
</tr>
<tr>
<td>Accuracy of volume reading</td>
<td>Estimated</td>
</tr>
<tr>
<td>Accuracy of Flow reading</td>
<td>+/-10%</td>
</tr>
<tr>
<td>Inspiratory Trigger</td>
<td>0.14 – 2.21 L/sec²</td>
</tr>
<tr>
<td>Expiratory Trigger</td>
<td>0.28 – 1.67 L/sec²</td>
</tr>
<tr>
<td>Low Flow Alarm</td>
<td>0 – 200 lpm</td>
</tr>
<tr>
<td>High Flow Alarm</td>
<td>0 – 200 lpm</td>
</tr>
<tr>
<td>Inspiratory Time</td>
<td>0.7 – 3.0 seconds</td>
</tr>
<tr>
<td>Back-up Rate</td>
<td>6 - 43 Breaths per minute</td>
</tr>
<tr>
<td>Type of protection against electric shock</td>
<td>Class 1 equipment</td>
</tr>
<tr>
<td>Degree of protection against electric shock</td>
<td>Type B to EN 60601-1</td>
</tr>
<tr>
<td>Mode of operation</td>
<td>Continuous</td>
</tr>
<tr>
<td>IP rating</td>
<td>X0</td>
</tr>
<tr>
<td>Storage environment</td>
<td>-20 to 50°C 5 – 85% RH 260 – 1100 mBar atmospheric pressure</td>
</tr>
<tr>
<td>Internal battery</td>
<td>18.75Vdc 116Whr</td>
</tr>
<tr>
<td>Running time</td>
<td>4 -12 hours depending on settings and leak</td>
</tr>
<tr>
<td>External battery</td>
<td>18.75Vdc 116Whr</td>
</tr>
<tr>
<td>Running time</td>
<td>4 -12 hours depending on settings and leak</td>
</tr>
</tbody>
</table>

**Protection against flammable anaesthetic mixtures** - Not suitable for use in the presence of a FLAMMABLE ANAESTHETIC MIXTURE WITH AIR OR WITH OXYGEN OR NITROUS OXIDE

**International Standards**
BS EN60601-1 1990, EN 10651- 6 2004
Safety of Electromedical Instruments, General Requirements
Electromagnetic Compatibility (In accordance with the EMC Directive 89/336/EMC)
Breas Medical Ltd declares that the NIPPY 3+ Ventilator complies with the following EMC standards. EN60601-1-2: 2001
Test results available for review from Breas Medical Ltd

[CE Mark: 0086]
Operation Under Extreme Conditions

Ambient Temperature in the range of +5 to +50 °C
Between 5 and 40 degrees functioning of the ventilator should not be affected. Extremes of temperature (below 5 °C, above 40 °C) may affect the colour of the LCD display. This will return to normal with the temperature. Operation above 40 degrees is not recommended. The ventilator may overheat at elevated temperatures. An audible and visual alarm will be activated in the event of over temperature. Air conditioning should be employed to keep the room temperature below 40 degrees.

Ambient Relative Humidity in the range of 10 to 100% RH
The ventilator is expected to function correctly at extremes of humidity. High humidity levels may affect the colour of the LCD display. This will return to normal with the humidity.

Atmospheric Pressure in the range of 600mBar to 1100mBar
The ventilator is expected to function correctly between 600 and 1100 mBar.

Supply Voltage Range from –20% to +10% of specified value
The NIPPY 3+ will operate normally

Failure of Electrical Power Supply
If a back-up battery is connected, the ventilator will automatically switch to the back-up supply and give an audible and visual indication that it has done so. During total power failure, there will be no output from the machine. The patient will be able to breathe spontaneously through the machine and out through the exhale port. However, some re-breathing of exhaled gas is inevitable. During power/ventilator failure disconnect the patient from the breathing circuit as soon as possible.

The inspiratory / expiratory resistance of the NIPPY 3+ and breathing system (NIPPY 3+ and circuit) is less than 6cm H₂O @ 60 l/min. This value must not be exceeded when adding attachments or fittings to the breathing circuit.

Accessories and Spares

- A range of nasal and facemasks is available in various sizes. Please contact us for details
- Head Set pt.no. 0563 available in Small, Medium, Large and Extra large. Please add S,M,L OR XL to part number when ordering.
- A range of breathing circuits is available for use with nasal mask, facemask or tracheotomy. These can be supplied with a heated wire for use with an external humidifier. See Breathing Circuits section.
- Air Filter Element pt.no. 0584 (pack of 5)
- Inline Bacterial Filter pt.no. 0635. - 99.999% filtration – Resistance, 0.75mB @ 50 l/min – deadspace 55ml – 22mm tapered fittings.

These components are for single patient use.
- External battery, part number 0910.
WARNINGS

This ventilator is intended to augment the patient breathing. It **MUST NOT BE USED AS A LIFE SUPPORT VENTILATOR.** It is not intended to provide the total ventilatory requirement of the patient.

Do not attempt to pass oxygen into the panel mounted air inlet, or use with flammable anaesthetic agents e.g. Ether etc.

The NIPPY 3+ must be connected to a grounded (earthed) electrical supply. The protective earth of the domiciliary electrical installation shall be checked for safe and effective operation.

CAUTIONS

- The NIPPY 3+ should only be used in accordance with the instructions of the supervising physician. **Personnel using and operating the NIPPY 3+ must become familiar with this instruction manual before using the unit.**

- Ensure patient safety through the presence of a trained attendant and an alternative means of ventilation. Consideration should also be given to the use of secondary alarm monitoring.

- The NIPPY 3+ should not be placed close to high frequency surgical diathermy, defibrillator or short wave therapy equipment as it may adversely effect the operation.

- The functioning of the ventilator can be adversely affected by electromagnetic interference exceeding the level of 10V/m in the test conditions of EN60601-1-2. E.g. Mobile telephone operation may adversely affect the operation of the ventilator.

- If the NIPPY 3+ is moved from cold surroundings into a well-heated room, condensation may form. Do not operate the unit for at least 2 hours to allow any condensation to evaporate.

- Do not operate the ventilator in direct sunlight.

- Avoid places where there is excessive humidity or dust, which may cause damage to internal parts.

- Keep the NIPPY 3+ away from extreme direct heat, such as fires, heating radiators etc., and always allow a 100mm (4.0in) air space around the unit when in use.

- If liquids are allowed to enter the unit, serious damage could occur. If you spill any liquid into the NIPPY 3+, consult qualified service personnel.

- Do not place any form of cover over the ventilator, especially near the air intake.

- DO NOT use anti static or electrically conductive tubing.

- Adding extra components / subassemblies to the breathing circuit may cause the pressure, during expiration, at the patient connection port of the breathing circuit to increase.
Using Supplementary Oxygen with the NIPPY 3+

If required, supplementary oxygen may be entrained into the breathing circuit up to a maximum of 15 L/minute.

When adding oxygen, fit an entrainment port at the mask / tracheotomy end of the circuit.

Switch on the NIPPY before the oxygen.

When treatment is complete, switch off and disconnect the oxygen supply. Switch off the NIPPY and disconnect the breathing circuit. Store the breathing circuit in a clean bag or other suitable container.

DO NOT leave the oxygen connected when not in use. This can cause a build-up of oxygen in, or around the machine.

DO NOT block the end of the breathing circuit with oxygen connected.

DO NOT expose oxygen to naked flames.

DO NOT smoke in the vicinity.

DO NOT use a gas cooker in the vicinity.

DO NOT use a gas, oil or solid fuel heater in the vicinity.

Precaution: always follow user instructions when entraining Oxygen.
MAINTENANCE

YOU MUST DISCONNECT THE NIPPY FROM THE MAINS SUPPLY BEFORE ANY MAINTENANCE IS CARRIED OUT

User maintenance is limited to cleaning and visual inspection of the ventilator, the input air filter and the breathing circuit.

The ventilator and the detachable mains cord set should be inspected for signs of external damage weekly. If any damage is evident (particularly to the mains cord set) refer repair to appropriately qualified technical personnel.

DO NOT immerse the ventilator in or spray with water

DO NOT use solvent cleaning agents or detergents

DO NOT use abrasive cleaning agents

**Mains Power Lead**
Before using the NIPPY, inspect the mains lead for damage. Do not use if there is any damage to the plug, socket or the insulation.

**Exterior of Case**
To clean, wipe the exterior of the case with a soft cloth moistened with water.

**Input Air Filter**
The input air filter should be inspected weekly. It is located on the rear of the machine. To remove the filter, grip the filter housing with the thumb and forefinger, across the bottom corners and pull the filter cover away from the ventilator. Remove and inspect the element. If the filter element requires replacement, use only recommended spares (see spares list). The use of any other filtering material may impair the performance of the ventilator.

Never attempt to clean the filter element with solvent cleaning agents.

**Do not** operate the ventilator unless the input air filter is in place.

**User Maintenance Schedule**

<table>
<thead>
<tr>
<th></th>
<th>Before Use</th>
<th>Daily</th>
<th>Weekly</th>
<th>Monthly</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alarms</td>
<td>Test</td>
<td>Test</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Batteries</td>
<td></td>
<td>Test</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Breathing Circuit</td>
<td>Inspect</td>
<td>Replace</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Inlet Filter</td>
<td>Inspect/Replace</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Power Cord</td>
<td>Inspect</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Battery Run Time Test**
Perform a run time test as detailed in the battery care section of this manual.

**Breathing Circuit Cleaning**
The breathing circuit is considered disposable.
Servicing / Repair

Only suitably qualified technically competent personnel should attempt servicing or repair of this ventilator.

To maintain its performance, the ventilator will require periodic servicing at the following intervals:
- Annual electrical safety test, alarm function, calibration and battery condition tests.
- 10000 hours use. The service reminder symbol will be displayed on screen.
Details of service requirements are contained in the technical manual.
Damage to either the machine or its mains lead must be inspected by competent technical personnel before use.
We offer various NIPPYcare service contracts for your ventilator. Please contact our sales team for further details of technical training or contracts: 01789 293460.

Technical Information

A technical manual incorporating circuit diagrams and descriptions will be made available, on request, to enable appropriately qualified technical personnel to repair the parts of the equipment designed to be repairable.

Warranty

The NIPPY 3+ is covered by a full 12 months parts and labour warranty, provided that the unit is properly operated under conditions of normal use. This warranty does not apply to any unit that has been subjected to misuse or accidental damage, or repaired or modified by unauthorised personnel.

Transportation

When shipping, damage as a result of inadequate packing is the customer's responsibility. Use the original packing materials whenever possible. Breas Medical Ltd are not responsible for inbound transit damage.

Factory Service / Repair

Breas Medical Ltd products returned for factory service or repair must have a Return Material Authorisation (RMA) number assigned. This is essential for efficient processing of repairs.
You can obtain your RMA number by calling 01789 293460 with the following information:
1. Unit Model
2. Serial number
3. Your name, address and telephone number
4. Complete description of the malfunction or service required

When the RMA number has been issued, we will arrange for the unit to be collected. Place the RMA number on the outside of the carton.
The unit must be properly packaged before shipment. Preferably, in the original packaging.
When enquiring about a returned item, you must quote the RMA number.

Disposal at end of Life

The NIPPY 3+ should be disposed of in line with local authority guidelines / regulations.
Spent batteries should be disposed of in line with local authority guidelines / regulations.
# EMC Information

**Guidance and Manufacturer's Declaration – Electromagnetic Emissions:**

This device is intended for use in the electromagnetic environment specified below. The user of this device should make sure it is used in such an environment.

<table>
<thead>
<tr>
<th>Emissions Test</th>
<th>Standard</th>
<th>Electromagnetic Environment- Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF emissions (radiated)</td>
<td>EN55011</td>
<td>The device uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.</td>
</tr>
<tr>
<td>CISPR 11</td>
<td></td>
<td></td>
</tr>
<tr>
<td>RF emissions (conducted)</td>
<td>EN55011</td>
<td>The device is suitable for use in all establishments, including domestic establishments and those directly connected to the public low-voltage power supply network.</td>
</tr>
<tr>
<td>CISPR 11</td>
<td>EN61000-3-2</td>
<td></td>
</tr>
<tr>
<td>Harmonic emissions</td>
<td>EN61000-3-2</td>
<td></td>
</tr>
<tr>
<td>IEC 61000-3-2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Voltage fluctuations/Flicker emissions</td>
<td>EN61000-3-2</td>
<td></td>
</tr>
<tr>
<td>IEC 61000-3-3</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

---

**Electromagnetic Immunity:**

This device is intended for use in the electromagnetic environment specified below. The user of this device should make sure it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity Test</th>
<th>IEC 60601 Test Level</th>
<th>Compliance Level</th>
<th>Electromagnetic Environment- Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrostatic Discharge (ESD)</td>
<td>±6 kV contact</td>
<td>±6 kV contact</td>
<td>Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.</td>
</tr>
<tr>
<td>IEC 61000-4-2</td>
<td>±8 kV air</td>
<td>±8 kV air</td>
<td>Mains power quality should be that of a typical home or hospital environment.</td>
</tr>
<tr>
<td>Electrical Fast Transient/Burst</td>
<td>±2 kV for power supply lines</td>
<td>±2 kV for supply mains</td>
<td>Mains power quality should be that of a typical home or hospital environment. If the user of the device requires continued operation during power mains interruptions, it is recommended that the device be powered from an uninterruptible power supply or a battery.</td>
</tr>
<tr>
<td>IEC 61000-4-4</td>
<td>±1 kV for input-output lines</td>
<td>±1 kV for input/output lines</td>
<td></td>
</tr>
<tr>
<td>Surge</td>
<td>±1 kV differential mode</td>
<td>±1 kV differential mode</td>
<td>Mains power quality should be that of a typical home or hospital environment.</td>
</tr>
<tr>
<td>IEC 61000-4-5</td>
<td>±2 kV common mode</td>
<td>±2 kV for common mode</td>
<td></td>
</tr>
<tr>
<td>Voltage dips, short interruptions and voltage variations on power supply input lines</td>
<td>&gt;95% dip in Voltage for 0.5 periods @ 230Vac and 100Vac</td>
<td>&gt;95% dip in Voltage for 0.5 periods @ 230Vac and 100Vac</td>
<td></td>
</tr>
<tr>
<td>IEC 61000-4-11</td>
<td>60% dip in Voltage for 5 periods @ 230Vac and 100Vac</td>
<td>60% dip in Voltage for 5 periods @ 230Vac and 100Vac</td>
<td></td>
</tr>
<tr>
<td>Power frequency magnetic field (50/60 Hz)</td>
<td>3 A/m</td>
<td>3 A/m</td>
<td>Power frequency magnetic fields should be at levels characteristic of a typical location in a typical hospital or home environment.</td>
</tr>
<tr>
<td>IEC 61000-4-8</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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Electromagnetic Immunity:

This device is intended for use in the electromagnetic environment specified below. The user of this device should make sure it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity Test</th>
<th>IEC 60601 Test Level</th>
<th>Compliance Level</th>
<th>Electromagnetic Environment Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conducted RF</td>
<td>3 Vrms</td>
<td>3 Vrms</td>
<td></td>
</tr>
<tr>
<td>IEC 61000-4-6</td>
<td>150 kHz to 80 MHz</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Radiated RF</td>
<td>3 V/m</td>
<td>3 V/m</td>
<td></td>
</tr>
<tr>
<td>IEC 61000-4-3</td>
<td>80 MHz to 2.5 GHz</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Recommended separation distances**:

- For conducted RF:
  - $150 \text{ kHz to } 80 \text{ MHz}$: $d = 1.2 \sqrt{P}$ @ $150 \text{ kHz to } 80 \text{ MHz}$
- For radiated RF:
  - $3 \text{ V/m}$ @ $80 \text{ MHz to } 2.5 \text{ GHz}$

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

**NOTE 1**: At $80 \text{ MHz}$ and $800 \text{ MHz}$, the higher frequency range applies.

**NOTE 2**: These guidelines may not apply in all situations. Electromagnetic propagation is affected by the proximity of structures, objects, and people.

a: Field strengths from transmitters, such as base stations for radio (mobile/cordless) telephones, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the device is used exceeds the applicable RF compliance level above, the device should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as relocating the device.

**Recommended Separation Distances between Portable and Mobile RF Communications Equipment and This Device**:

The device is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The user of this device can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and this device as recommended below, according to the maximum output power of the communications equipment.

<table>
<thead>
<tr>
<th>Rated Maximum Power Output of Transmitter (W)</th>
<th>Separation Distance According to Frequency of Transmitter (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>150 kHz to 80 MHz</td>
<td>80 MHz to 800 MHz</td>
</tr>
<tr>
<td>$d = 1.2 \sqrt{P}$</td>
<td>$d = 1.2 \sqrt{P}$</td>
</tr>
<tr>
<td>0.01</td>
<td>0.12</td>
</tr>
<tr>
<td>0.1</td>
<td>0.38</td>
</tr>
<tr>
<td>1</td>
<td>1.2</td>
</tr>
<tr>
<td>10</td>
<td>3.8</td>
</tr>
<tr>
<td>100</td>
<td>12</td>
</tr>
</tbody>
</table>

For transmitters rated at a maximum output power not listed above, the recommended separation distance $d$ in meters (m) can be estimated using the equation applicable to the frequency of the transmitter, where $P$ is the maximum output power rating of the transmitter in watts (W).

**Note 1**: At $80 \text{ MHz}$ and $800 \text{ MHz}$, the separation distance for the higher frequency range applies.

**Note 2**: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects, and people.
## Typical Power Output of Some Common Transmitters

This list is provided for general guidance. It is not exhaustive or specific. It is not intended to replace the findings of an electromagnetic survey.

<table>
<thead>
<tr>
<th>Power</th>
<th>Notes</th>
<th>Suggested Minimum Separation Distance</th>
</tr>
</thead>
<tbody>
<tr>
<td>100 kW</td>
<td>Typical transmission power of FM radio station with 50 km range</td>
<td>727m</td>
</tr>
<tr>
<td>1 kW = 1000 W</td>
<td>Maximum allowed output RF power from a amateur radio transceiver without special permissions</td>
<td>73m</td>
</tr>
<tr>
<td>100 W</td>
<td>Typical maximum output RF power from an amateur radio transceiver</td>
<td>23m</td>
</tr>
<tr>
<td>5 W</td>
<td>Typical maximum output RF power from a hand held amateur radio transceiver</td>
<td>5m</td>
</tr>
<tr>
<td>4 W</td>
<td>Typical maximum output power for a Citizens’ band radio station, (27 MHz) in many countries</td>
<td>4.6m</td>
</tr>
<tr>
<td>2 W</td>
<td>Maximum output from a UMTS/3G mobile phone (Power class 1 mobiles)</td>
<td>3.25m</td>
</tr>
<tr>
<td>500 mW</td>
<td>Maximum output from a UMTS/3G mobile phone (Power class 2 mobiles)</td>
<td>1.6m</td>
</tr>
<tr>
<td>400 mW</td>
<td>Access point for Wireless networking</td>
<td>1m</td>
</tr>
<tr>
<td>250 mW</td>
<td>Maximum output from a UMTS/3G mobile phone (Power class 3 mobiles)</td>
<td>1.15m</td>
</tr>
<tr>
<td>32 mW</td>
<td>Typical WiFi transmission power in laptops.</td>
<td>400cm</td>
</tr>
<tr>
<td>2.5 mW</td>
<td>Bluetooth Class 2 radio, 10 m range</td>
<td>115cm</td>
</tr>
<tr>
<td>1.0 mW = 1000 µW</td>
<td>Bluetooth standard (Class 3) radio, 1 m range</td>
<td>7.2cm</td>
</tr>
<tr>
<td>100 µW</td>
<td>Typical maximum received signal power (−10 to −30 dBm) of wireless network</td>
<td>2.3cm</td>
</tr>
</tbody>
</table>
NIPPY+ Battery run time test record

Use this table to record the battery run times when performing the monthly battery health test.

Replace the battery when the run time is less than 75% of a new battery or the battery is 2 Years old.

Fully charge the battery, connect the breathing circuit and using your normal settings run the ventilator until the low battery alarm operates. Record the run time in the table.

NIPPY Serial Number:…………………………..

<table>
<thead>
<tr>
<th>Date</th>
<th>Runtime – Internal battery</th>
<th>Runtime – External Battery If applicable</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
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</tr>
</tbody>
</table>
Locking the settings

Adjustment lock
The settings can be locked to prevent unauthorised adjustment.

To lock press ◀- and ◀+ buttons simultaneously and hold for 2 seconds.

To unlock press ◀- and ◀+ buttons simultaneously and hold for 2 seconds.

This prevents adjustment but allows the user to switch the ventilator on and off.

Total lock
The ventilator can be locked to prevent unauthorised adjustment or power off.

To lock press ◀- and ◀+ and SET buttons simultaneously and hold for 2 seconds.

To unlock press ◀- and ◀+ and SET buttons simultaneously and hold for 2 seconds.

In this mode, the ventilator must be unlocked before it can be switched off.

This page may be removed before the instruction manual is passed to the user.