

USER MANUAL





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Masimo SET is the pulse oximetry technology proven to provide accurate measurements under the challenging monitoring conditions of low perfusion and patient motion as supported by over 100 clinical studies.

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Using any sensors other than Nonin-branded PureLight® sensors with the Nonin Xpod USB connector may result in inaccurate performance (of the Airvo™ 3 and/or Nonin products) and will void the Nonin product warranty.

Before you start

- This user manual is for instructions on using the Airvo 3 NIV.
- This user manual is intended for healthcare professionals. While the information provided is believed to be accurate, it is not a substitute for exercising professional judgement.
- Read this user manual, including all warnings, before using the Airvo 3 NIV.
- Before the Airvo 3 NIV is used for the first time, it must be set up according to the instructions in the Airvo 3 Technical Manual.
- Some accessories may not be available in certain countries. Please contact your local Fisher & Paykel Healthcare representative for more information.
- If any device or accessory label is damaged or unreadable, contact your Fisher & Paykel Healthcare representative for a replacement.

Additional resources

- If using the Disinfection Kit to reprocess the Airvo 3 NIV, refer to the Disinfection Kit Manual provided with the Disinfection Kit (900PT600).
- Refer to user instructions supplied with individual accessories for correct use and additional safety information.
- Refer to the Airvo 3 Technical Manual for initial setup, maintenance, servicing and additional troubleshooting instructions.
- Visit the Airvo 3 website at: www.fphcare.com/airvo3 to download user instructions including this user manual.
- For assistance from your Fisher & Paykel Healthcare representative contact us at: www.fphcare.com/contact-us.

Conventions used in this manual

Warning

A warning alerts the user to a potential hazard with use or misuse of the device which, if not avoided, could result in death or serious injury.

Caution

A caution alerts the user to a potential hazard with use or misuse of the device which, if not avoided, could result in minor or moderate injury.

Note

A note emphasizes information important for using the Airvo 3 NIV correctly.

Contents

Before you start	1
1 Introduction	4
1.1 Intended/indications for use.....	4
1.2 Contraindications	4
1.3 Side-effects	4
2 Safety information	4
2.1 General.....	4
2.2 Supplementary oxygen	6
3 Overview	7
3.1 Identifying system components.....	7
3.2 Identifying device components	8
3.3 Navigating the user interface	9
4 Preparing the Airvo 3 NIV	11
4.1 Equipment required	11
4.2 Airvo 3 NIV setup	13
4.3 Additional setup for adult NIV	15
4.4 Supplementary oxygen	16
5 Using the Airvo 3 NIV	17
5.1 Getting started.....	17
5.2 Selecting the therapy	19
5.3 Optiflow high flow therapy settings.....	20
5.4 Starting Optiflow high flow therapy	21
5.5 Adult NIV therapy settings	24
5.6 Starting adult NIV therapy.....	25
5.7 During therapy	28
5.8 Mobility and battery operation.....	29
5.9 Changing therapy	30
5.10 Stopping therapy.....	31
6 Monitoring data	32
6.1 Patient data.....	32

6.2	Live value graphs	33
6.3	Long term graphs	34
7	Troubleshooting	35
7.1	Alarms	35
7.2	Alarm priority	35
7.3	Auditory information signals	35
7.4	Viewing alarm details	36
7.5	Checking the alarm system	36
7.6	Airvo 3 NIV alarms	36
8	Reprocessing	42
8.1	Airvo 3 NIV device exterior reprocessing	42
8.2	Outlet elbow reprocessing	43
8.3	Schedule for changing accessories	45
8.4	Replacing the air filter	46
8.5	Servicing	46
9	Pulse oximetry	46
9.1	Pulse oximetry warnings, cautions, and notes	46
9.2	Setup for pulse oximetry	47
9.3	During therapy	48
9.4	Description of measurements	50
9.5	Description of setting and alarms	51
9.6	Alarm and measurement settings	52
9.7	Troubleshooting	53
10	Adult NIV therapy	55
10.1	Therapy description	55
10.2	Alarm settings	61
	Specifications	62
	Glossary	68
	Appendix 1. Patient consumables	69
	Appendix 2. Parts and accessories	72
	Appendix 3. Pulse oximetry accessories	73
	Appendix 4. Humidification behavior during battery operation	77

1. Introduction

The Airvo 3 NIV is designed to deliver:

- Optiflow™ high flow, and
- noninvasive ventilation (NIV)

to spontaneously breathing patients.

A blower inside the Airvo 3 NIV entrains room air, which may be blended with oxygen from high-pressure sources (such as wall supplies or bottles) or low-pressure sources (such as flowmeters). The air-oxygen mixture is then warmed and humidified in the water chamber, before being transported through the heated breathing tube to the patient interface.

The Airvo 3 NIV is powered by wall power supply, with internal battery backup to provide continuity of therapy during intra-hospital transport.

1.1 Intended use/indications for use

When used in NIV mode:

The Airvo 3 NIV provides non-invasive ventilatory support for non-ventilator-dependent, spontaneously breathing adult patients (30 kg and above) with respiratory insufficiency, or respiratory failure. It is intended to be used in hospital facilities. It is not intended for life support.

When used in High Flow mode:

The Airvo 3 NIV is for the treatment of spontaneously breathing patients who would benefit from receiving high flow warmed and humidified respiratory gases. This includes patients who have had upper airways bypassed. The flow may be from 2 – 70 L/min depending on the patient interface. The Airvo 3 NIV is for patients in hospital facilities.

The Airvo 3 NIV can deliver these high flow gases through nasal cannula to augment the breathing of spontaneously breathing neonate, infant, child, adolescent and adult patients suffering from respiratory distress and/or hypoxemia in the hospital setting. The Airvo 3 NIV is not intended to provide total ventilatory requirements of the patient and is not for use during field transport.

1.2 Contraindications

Contraindications are therapy-specific. Refer to instructions of patient interfaces and/or tube and chamber kits for therapy-specific contraindications.

1.3 Side-effects

The following side effects may occur during NIV therapy:

- Nasal, mouth or throat dryness
- Nosebleeds
- Abdominal bloating
- Ear or sinus discomfort
- Eye irritation
- Aspiration of fluid

Side effects are therapy-specific. Refer to instructions of patient interfaces and/or tube and chamber kits for therapy-specific side-effects.

2. Safety information

The Airvo 3 NIV and accessories are to be operated by, or under the supervision of, qualified personnel only. Read this manual, particularly all warnings, cautions and notes, and the instructions for use supplied with all accessories before using the Airvo 3 NIV.

2.1 General

Warnings

- The Airvo 3 NIV is not intended for life support. Do not use Airvo 3 NIV on patients who cannot tolerate a brief interruption of therapy.
- Appropriate patient monitoring is required for all patients using the Airvo 3 NIV.
- Delivery of respiratory gases may generate positive airway pressure. This must be considered where positive airway pressure could have adverse effects on a patient. To avoid serious injury, appropriately monitor the patient for risk factors of airway and lung pressure injury.

- Anybody connecting patient consumables, accessories or spare parts to the Airvo 3 NIV is accountable for the compatibility of the device and those patient consumables, accessories and/or spare parts.
- Do not use any patient consumables, accessories or spare parts that are not listed in this user manual, or the Airvo 3 Technical Manual. Incompatible consumables, parts or accessories could affect the quality of therapy, injure the patient, decrease electromagnetic immunity or increase electromagnetic emissions.
- Use only patient interfaces, heated breathing tubes, water chambers and filters specified in this manual to prevent disconnection during use, especially when moving the Airvo 3 NIV.
- Do not use antistatic or electrically conductive patient breathing circuits with the Airvo 3.
- Do not connect the Airvo 3 NIV to the battery of a battery-powered wheelchair, which may compromise device performance and therapy delivered.
- Do not start or operate the Airvo 3 NIV unless the device setup, including all accessories, is verified to be correct.
- Carefully route accessories, cords and cables, including the breathing tube, to reduce the possibility of patient entanglement or strangulation.
- Visually inspect the Airvo 3 NIV and accessories before use and replace if damaged or suspected to be damaged. Using a damaged device or accessories may impair performance and/or compromise safety.
- Make sure the auditory alarm signal is audible to the operator who will respond to alarms by following the instructions in section 7.5 to test the alarm before starting therapy.
- Do not use an Airvo 3 NIV on more than one patient at any one time.
- Do not use accessories beyond the maximum period of use specified in this manual. Exceeding the maximum use period can result in serious injury, including infection.
- Do not expose the Airvo 3 NIV battery to water, fire or excessive heat. Do not crush, disassemble or puncture the battery, or short-circuit the connector terminals.
- Do not use the Airvo 3 NIV as an apnea monitor.
- Do not use the Airvo 3 NIV for arrhythmia analysis.
- In the event of a battery leaking, do not allow the liquid to come in contact with the skin or eyes. If contact has been made, wash the affected area with copious amounts of water and seek medical advice.
- Seek medical advice immediately if a cell or a battery has been swallowed.
- Changes or modifications not expressly approved by Fisher & Paykel Healthcare voids the user's authority to operate the device.
- The therapy delivered to the patient can be impacted by the use of a pneumatic/jet nebuliser. Refer to compatible accessories and drug manufacturer instructions for correct usage.
- Do not use any solutions, suspensions, emulsions, anesthetic or respirable gases that are not identified in these user instructions. They may not be compatible with the patient consumables, device or accessories.
- Use only genuine F&P replacement battery modules to prevent damage to the Airvo 3 NIV, including excessive temperatures, fire or explosion.

Operating environment

- Do not use the Airvo 3 NIV at an altitude or temperature outside the rated range listed in the specifications section of the manual. Using outside of these ranges can compromise the equipment performance which consequently can result in degradation of the health of the patient.
- Do not use the Airvo 3 when outside the operating conditions listed in the specifications section. Therapy may be compromised outside this range.
- Do not use the Airvo 3 NIV in a magnetic resonance imaging (MRI) environment.
- Do not use the Airvo 3 NIV with, or in the presence of, a flammable anesthetic mixture with air or oxygen.
- Do not use the Airvo 3 NIV, or accessories, during defibrillation.
- Do not use the Airvo 3 NIV, or accessories, near any ignition source, including electrosurgery, electrocautery, or laser surgery instruments. Exposure to oxygen increases the risk of fire that may result in patient injury.
- Do not use the Airvo 3 NIV, or accessories during electrocautery.
- Explosion hazard: Do not use the Airvo 3 NIV in the presence of flammable anesthetics or other flammable substance in combination with air, oxygen-enriched environments, or nitrous oxide.
- Do not use the Airvo 3 NIV in a hyperbaric chamber.
- Avoid using the Airvo 3 NIV, or accessories, adjacent to, or stacked with, other equipment, which could result in improper operation. If such use is necessary, this equipment and the other equipment should be observed to verify that they are operating normally.
- Do not use the Airvo 3 NIV when the patient is in an incubator or infant warmer.
- The Airvo 3 NIV is not designed for use in the home.

Caution

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

To avoid burns

- Do not touch the hot surface of the heater-plate or chamber base.
- Never operate the Airvo 3 NIV if:
 - the heated breathing tube has been damaged in any way including holes, tears or kinks,
 - it is not working properly, or
 - water has entered the device.
- Do not cover the ventilator or place in a position that affects proper operation.
- Do not block the flow of air through the Airvo 3 NIV or breathing tube.

To avoid electric shock

- Do not store or use the Airvo 3 NIV where it can fall, or be pulled, into water. Disconnect the power cord and stop using the Airvo 3 NIV if water has entered the case.
- To protect from electric shock, always remove the sensor and completely disconnect the pulse oximeter before bathing the patient.
- Never operate the Airvo 3 NIV if it has, or is suspected of having:
 - been dropped or damaged,
 - a damaged power cord or plug, or
 - been dropped into water.
- See the Airvo 3 NIV Technical Manual for instructions to replace a damaged power cord.
- Do not attempt to adjust, repair, open, disassemble or modify the Airvo 3 NIV, pulse oximeter equipment, or accessories, except as described in this user manual or the Airvo 3 Technical Manual. Return the Airvo 3 NIV to your Fisher & Paykel Healthcare representative for servicing, if necessary.
- Do not touch the patient at the same time as any conductive parts of the device, such as USB ports.

! Notes

- If a serious incident has occurred while using this device please inform your local Fisher & Paykel Healthcare representative and Competent Authority in your country.

2.2 Supplementary oxygen**! Warnings**

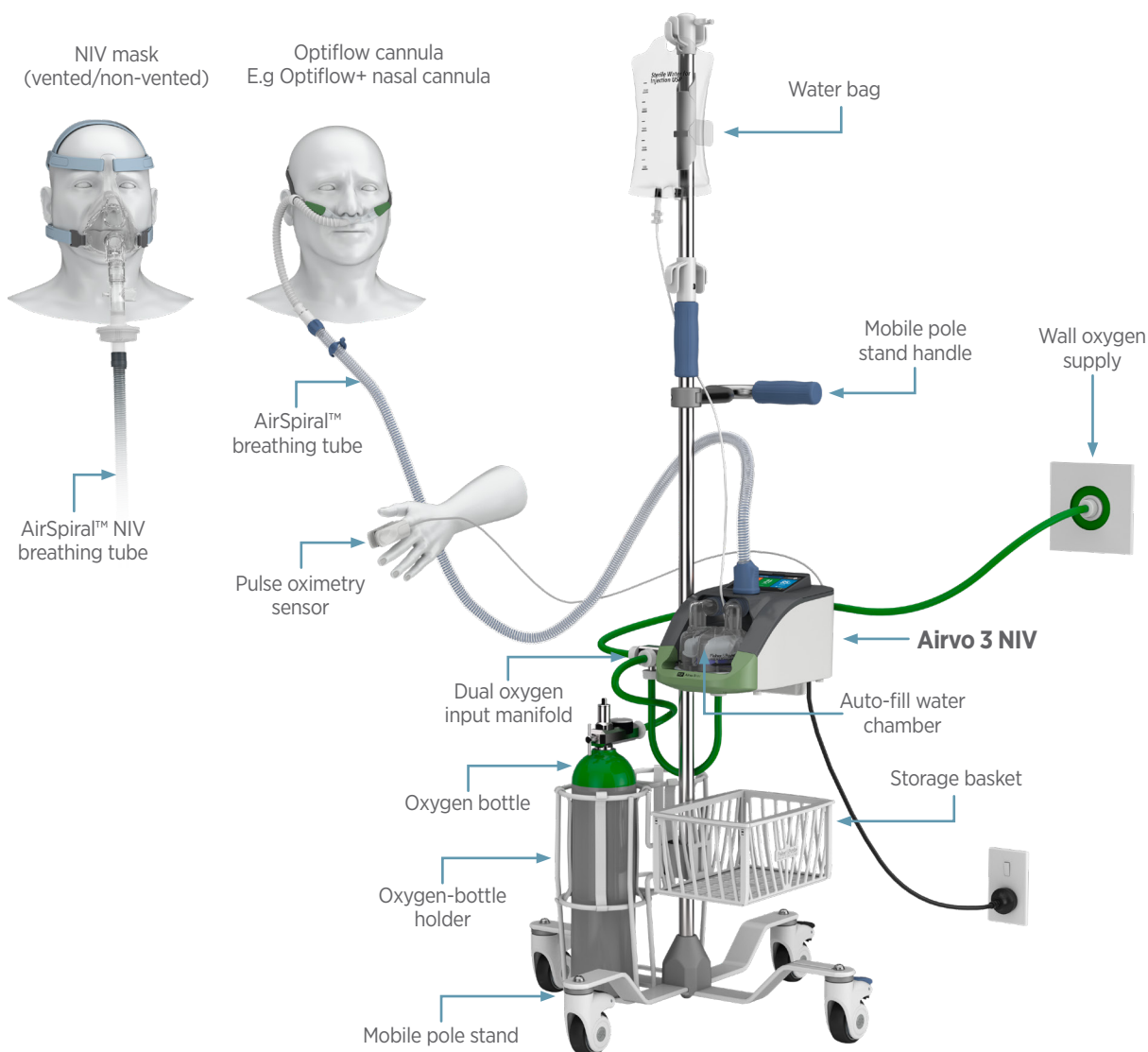
- You must take special care when using supplementary oxygen to reduce the risk of fire. Keep all sources of ignition away from the Airvo 3 NIV and, preferably, not in the same room as the Airvo 3 NIV during use.
- Do not use supplementary oxygen while smoking, near sparks or open flames.
- A spontaneous and violent ignition may occur if oil, grease or greasy substances contact oxygen under pressure. Keep these substances away from all oxygen equipment.
- The Airvo 3 NIV is a high flow device. Ensure the oxygen supply is designed to provide enough oxygen flow for all connected equipment, particularly when the supply is shared by multiple devices.
- Only connect pure oxygen to the oxygen inlet ports on the Airvo 3 NIV. The oxygen concentration displayed will be wrong if any other gas, or mixtures of gases, is connected.
- Only use lotions and/or salves that are labeled as oxygen-compatible to avoid the risk of fire and burns.

3. Overview

This section shows the Airvo 3 NIV system and compatible accessories.

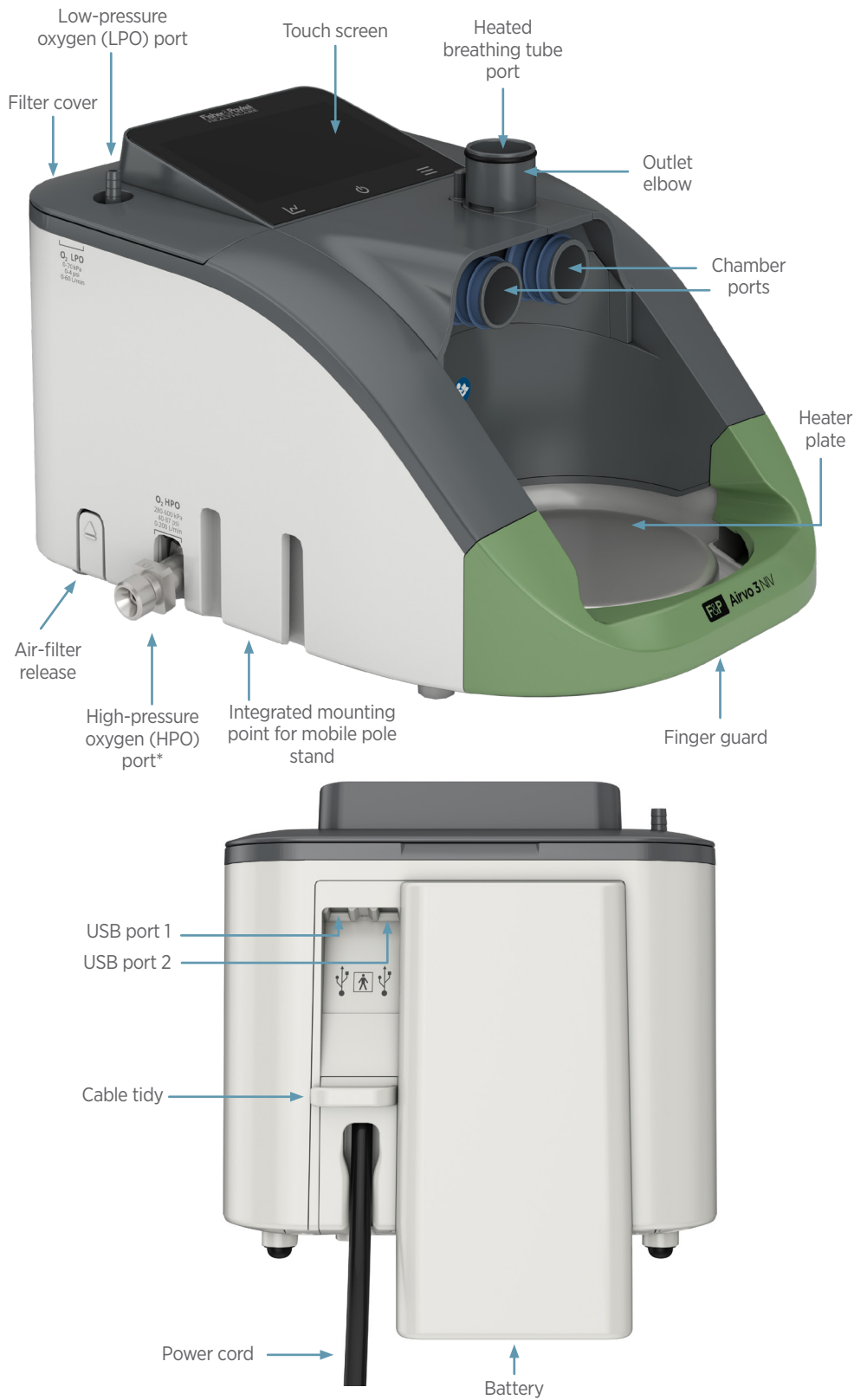
3.1 Identifying system components

Patient interface

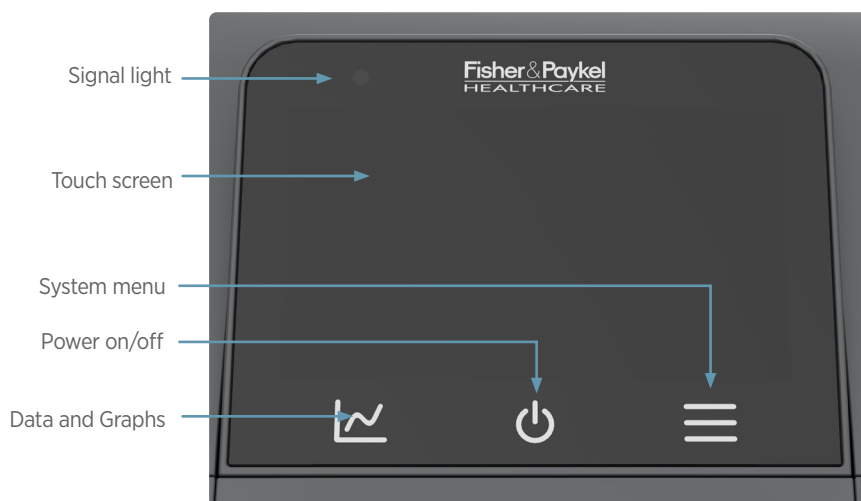


The Airvo 3 NIV System

3.2 Identifying device components



*HPO connection may vary depending on regional selection of connector type (DISS, NIST or SIS)



3.3 Navigating the user interface

The Airvo 3 NIV touch screen provides access to therapy and device status, settings and alarms. You interact with the user interface by:

- touching elements on the screen to open setting screens, make selections and change values, and
- swiping up/down to scroll through menus that are only partly displayed.

3.3.1 Home screen

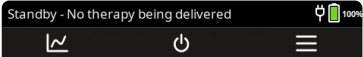


Warning

To ensure responsiveness keep Airvo 3 NIV touchscreen clean and dry. Performance may be reduced if the screen is allowed to become wet.

3.3.2 Message bar

The Message bar shows the current state of therapy delivery, confirms settings changes and displays alarms. Example messages are shown in the table below.

Message bar	Description
	Breathing gases are not being delivered to the patient. Tap the Start button to begin therapy.
	Breathing gases are being delivered. Tap the Stop button, then confirm action to return to standby mode.
	Active alarms are displayed on top of other messages. Tap the alarm for details or press  to temporarily pause the alarm audio. See section 7 for troubleshooting alarms.

3.3.3 Status indicators

The following icons may be displayed in the Message bar.

Icon	Description
	Audio pause
	The Airvo 3 NIV is being powered from the wall power supply
	Status of the internal battery
	50% of the battery charge is remaining
	Battery is charging and 50% of the charge is remaining
	Battery is not charging properly*
	Battery is missing or faulty*
	Battery is due for replacement*
	Touch display is locked to prevent accidental changes
	An Airvo 3 NIV USB storage device is connected to one of the USB ports

*Check the battery is properly installed. Replace the battery if the problem persists.

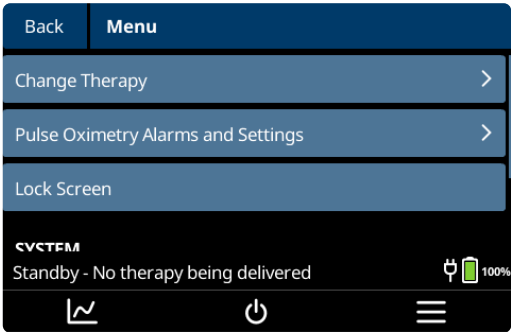
3.3.4 Signal light

The signal light flashes when any alarm is active. Its color indicates the highest priority alarm that is active. See section 7 for troubleshooting alarms.



3.3.5 System menu

The system menu provides access to additional settings and information. Tap  to open the system menu when the Home screen is displayed.



Menu item	Description
Change Therapy	Open the therapy selection screen to change between Optiflow high flow and NIV
NIV Advanced Settings	Change advanced settings for NIV therapy*
NIV Alarm Settings	Change settings for NIV alarms*
Pulse Oximeter Alarms and Settings	Configure the pulse oximetry settings including SpO ₂ alarms
Lock Screen	The lock screen can prevent accidental settings changes
Device Info	Displays the version, disinfection, and battery information
System Settings	Change advanced Airvo 3 NIV settings, limits and behaviors. Refer to the Airvo 3 Technical Manual for more information

* Only available when NIV therapy is active.

3.3.6 Data and Graphs screen

The Data and Graphs screen displays current and previous measurements and settings for the current patient.

Tap  to open the Data and Graphs screen when the Home screen is displayed. The values available depends on the active therapy mode.

4. Preparing the Airvo 3 NIV

Review the safety information in section 2 before proceeding. Refer to appendices 1 – 3 for a list of consumables and accessories that have been validated for use with the Airvo 3 NIV.

4.1 Equipment required

You will need:

- Airvo 3 NIV attached to a mobile pole stand,
- clean and disinfected outlet elbow,
- bag of USP sterile/distilled water for inhalation (or equivalent).

Outlet elbows can be processed in two different ways:

Disinfection kit (900PT600)

For hospitals using the disinfection kit for reprocessing: A clean and disinfected outlet elbow will already be installed in the Airvo 3 NIV. Remove the clean storage cover and/or the red disinfection tube before use.

Washer-disinfector

For hospitals using a washer-disinfector for reprocessing: obtain a clean and disinfected outlet elbow, e.g. from your Central Sterile Services Department (CSSD) system.

If supplementary oxygen is prescribed for your patient, you will need either:

- a high-pressure oxygen hoses to connect the Airvo 3 to the wall oxygen supply or an oxygen-bottle regulator, or
- low-pressure oxygen tubing to connect the Airvo 3 to a flowmeter.

Warnings

Only use patient consumables and accessories that are compatible with the Airvo 3 NIV (see Appendix 1-3).

Do not modify patient consumables or accessories in any way.

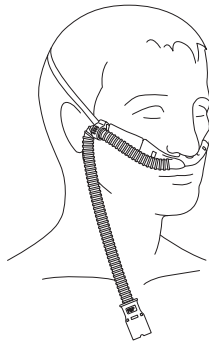
4.1.1 Optiflow high flow therapy

To provide Optiflow high flow therapy, you will need a:

1. Breathing tube and chamber kit.
2. Optiflow patient interface.

Refer to Appendix 1 for a list of compatible consumables.

Nasal interface

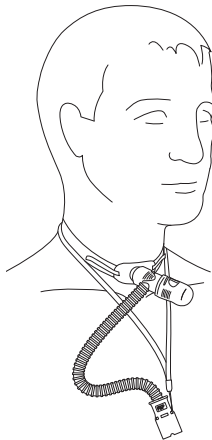


Optiflow+
Optiflow 3S
Optiflow+ Duet



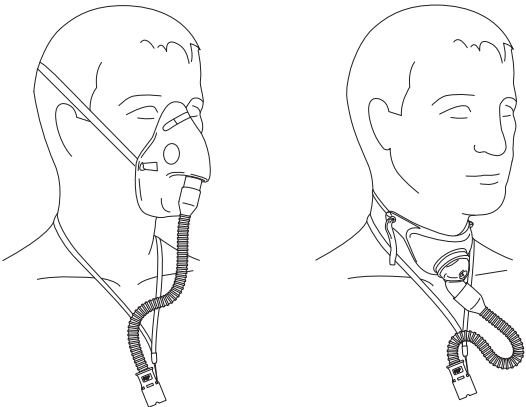
Optiflow Junior 2
Optiflow Junior 2+

Tracheostomy interface



Optiflow+ tracheostomy interface

Mask interface adapter



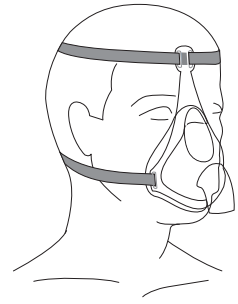
Optiflow+ mask interface adapter

4.1.2 Adult NIV

To provide adult NIV therapy, you will need a:

1. Vented or a non-vented NIV mask (with exhalation port) See Appendix 1 for compatible F&P patient interfaces
2. and either an:
 - AirSpiral NIV tube and chamber kit to set up adult NIV therapy for a new patient, or
 - AirSpiral NIV tube kit to change a patient's therapy from Optiflow high flow to adult NIV, reusing the patient's water chamber.

NIV interface



Vented/non-vented mask

Warnings

If using an oronasal or full face mask, the mask needs to have an anti-asphyxia valve.

When using a non-vented mask always use an exhalation port.

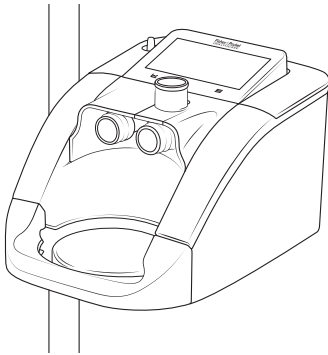
Only use the 900PT56X breathing tube kits and Optiflow interfaces when in Optiflow mode.

Only use the 900PT57X NIV breathing tube kits and NIV patient interfaces when in NIV mode.

In patients for whom CO₂ rebreathing is a significant concern, use a CO₂ monitor conforming with ISO 80601-2-55 to ensure CO₂ rebreathing is minimized.

4.2 Airvo 3 NIV setup

Standard aseptic techniques should be followed to minimize contamination when handling the Airvo 3 and accessories.



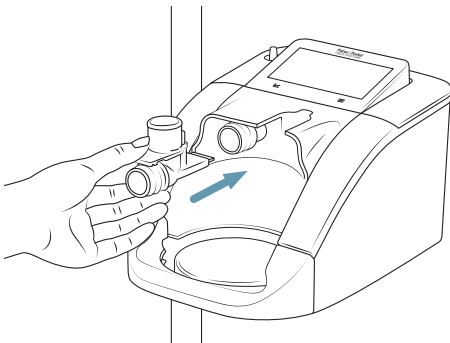
1. Check Airvo 3 NIV height

Check that the Airvo 3 NIV is attached securely to the mobile pole stand and is below the patient's head height.

Position the Airvo 3 NIV so that the power cord connection to the wall power supply is easily accessible and can be disconnected if necessary.

Caution

Do not place the Airvo 3 NIV where controls can be changed by the patient.



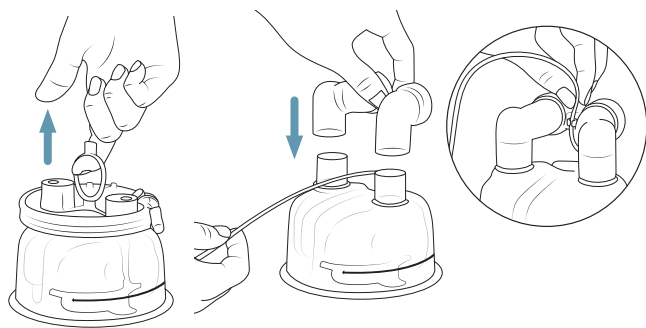
2. Connect the outlet elbow (if applicable)

This step applies if your hospital uses a washer-disinfector to clean and disinfect the outlet elbow. This step does not apply if your hospital uses the disinfection kit (900PT600).

Insert the clean, disinfected outlet elbow into the slot on the top of the Airvo 3 NIV.

Warning

Make sure the Airvo 3 NIV is off when connecting the outlet elbow.

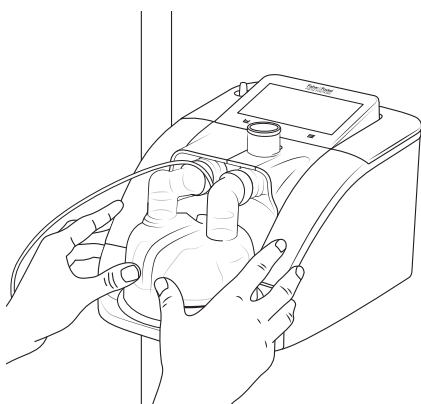


3. Assemble the water chamber

Open the tube and chamber kit and remove the MR290 auto-fill water chamber and chamber adapter.

Remove the blue port caps from the chamber by pulling the tear tab upwards then remove the bracket holding the water supply tube.

Fit the supplied adapter over the two vertical ports on the chamber and push on fully then clip the water supply tube into position.



4. Insert the water chamber

Fit the water chamber to the Airvo 3 NIV, sliding the chamber past the finger guard onto the heater-plate. Take care to align the port adapter with the blue ports on the Airvo 3 NIV.

Ensure the water chamber is fully inserted by pushing firmly on the front of the chamber until it slides past the finger guard.

To remove the water chamber, grip the port adapter and pull the chamber away from the Airvo 3 NIV.

Warnings

To avoid burns:

Do not start therapy without the water chamber in place.

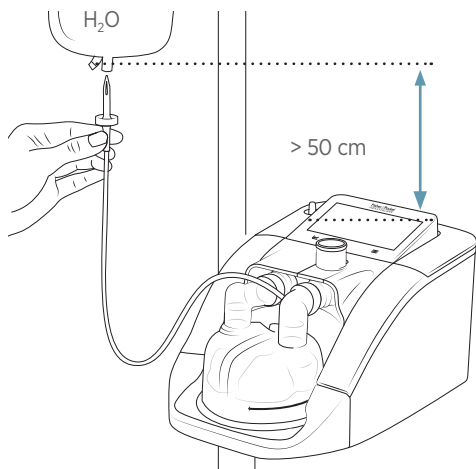
Do not touch the heater-plate, water chamber or chamber base during use.

Exercise caution when removing the chamber.

To avoid electrical shock:

When handling the Airvo 3 NIV with the water chamber in place, avoid tilting the device to prevent any chance of water entering the unit enclosure

Do not use the MR290 auto-fill water chamber if it has been dropped, allowed to run dry or damaged in any way. This could lead to the chamber overfilling.



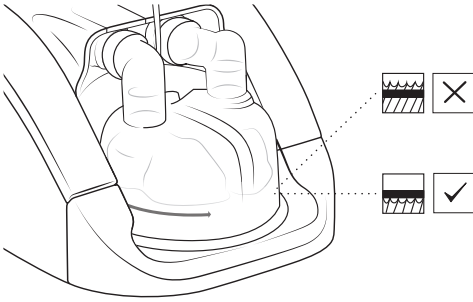
5. Connect the water bag

Attach the sterile water bag to the hanging bracket 50 cm above the Airvo 3 NIV. Remove the spike from the chamber bracket and push the bag spike into the fitting at the bottom of the bag.

Open the vent cap on the side of the bag spike.

Caution

Only use USP sterile/distilled water, suitable for inhalation, to fill the water chamber. Adding other substances can adversely affect the humidifier and therapy delivered.



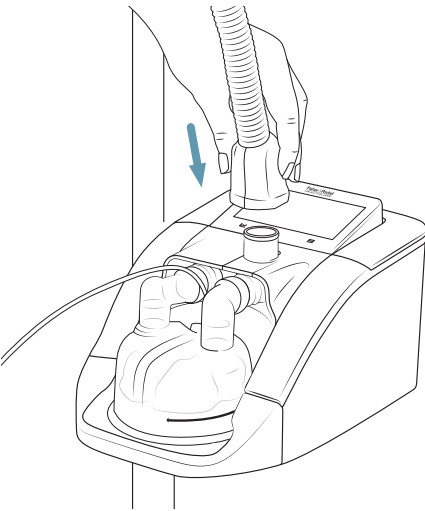
6. Check the water level

Check that water flows into the chamber and remains below the maximum water-level line.

The chamber will automatically maintain the correct water level until the water bag is empty.

Caution

Do not use the MR290 auto-fill chamber if the water level rises above the maximum water-level line. This may lead to water entering the patient's airway.



7. Install the breathing tube

Connect the breathing tube by lining up the pins on top of the Airvo 3 NIV, pushing down until you hear a click and the tube locks into place.

To remove the breathing tube, squeeze the sides of the connector and pull up.

Warning

To avoid burns:

Do not use an insulating sleeve or any similar accessories which are not recommended by Fisher & Paykel Healthcare.

Note

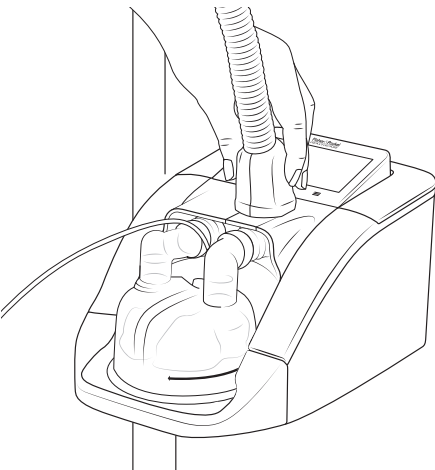
Make sure the outlet elbow is installed in the Airvo 3 NIV before attaching the heated breathing tube.

See step 2 "Connect the outlet elbow (if applicable)" above.

This step applies if your hospital uses a washer-disinfector to clean and disinfect the outlet elbow.

This step does not apply if your hospital uses the disinfection kit (900PT600).

Insert the clean, disinfected outlet elbow into the slot on top of the Airvo 3 NIV.



4.3 Additional setup for adult NIV

Follow the steps in the user instructions for the AirSpiral NIV kits to set up the Airvo 3 NIV system to deliver NIV therapy.

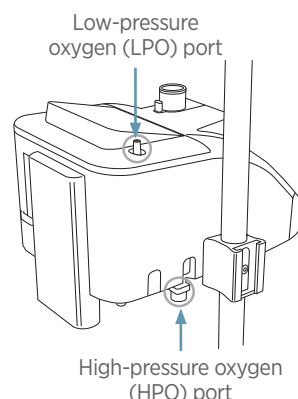
4.4 Supplementary oxygen

The Airvo 3 NIV provides two options for connecting supplementary oxygen

1. A high-pressure oxygen (HPO) inlet port, and
2. A low-pressure oxygen (LPO) inlet port.

The high-pressure oxygen inlet port is connected to the wall oxygen supply or to the pressure regulator on an oxygen bottle. The ability of the Airvo 3 NIV to provide the target FiO_2 is limited by the line pressure of the high-pressure inlet port (HPO). If the Airvo 3 NIV is unable to maintain the target FiO_2 , the device will generate a “ FiO_2 Below Target” alarm.

The low-pressure oxygen inlet port is connected to an external flowmeter, typically a rotameter.



Warnings

You must take special care when using supplementary oxygen to reduce the risk of fire. Keep all sources of ignition away from the Airvo 3 NIV and, preferably, not in the same room as the Airvo 3 NIV during use.

Do not use supplementary oxygen while smoking, near sparks or open flames.

When using bottled oxygen, ensure the volume remaining in the bottle is sufficient for the therapy planned.

Connect only pure oxygen gas to the oxygen inlet ports on the Airvo 3 NIV. The oxygen concentration displayed will be wrong if any other gas, or mixtures of gases, is connected.

The oxygen concentration delivered to the patient can be affected by changes to the oxygen setting, patient interface or obstructions in the air path.

Only use lotions and/or salves that are labeled as oxygen-compatible to avoid the risk of fire and burns.

Appropriate patient monitoring must be used at all times.

Make sure that all oxygen connectors are tightened sufficiently to prevent leaks.

As the low-pressure oxygen (LPO) inlet port uses an alternative small-bore connector design different from those specified in the ISO 80369 series, there is a possibility that a misconnection can occur with a medical device using a different alternative small-bore connector, which can result in a hazardous situation causing harm to the patient. Special measures need be taken by the user to mitigate these reasonably foreseeable risks.

During Optiflow high flow therapy, the fraction of oxygen inspired by the patient will be lower than the value displayed on the FiO_2 tile if the patient's peak inspiratory demand exceeds the flow delivered.

The Airvo 3 NIV is a high-flow device and should only be connected to a pipeline installation designed using a diversity factor that allows for the indicated high-flow at a specified number of terminal outlets, in order to avoid exceeding the pipeline design flow, thereby minimizing the risk that the humidifier interferes with the operation of adjacent equipment.

There is a risk of fire associated with oxygen enrichment during oxygen therapy. Do not use the equipment or accessories near sparks or open flames.

Smoking during oxygen therapy is dangerous and is likely to result in facial burns or death. Do not allow smoking or open flames within the same room as the equipment or any oxygen-carrying accessories. If the patient intends to smoke, always turn the equipment off, remove the cannula and leave the room where the equipment is located. If unable to leave the room, wait 10 minutes after the equipment has been turned off.

Do not lubricate fittings, connections, tubing, or other accessories of the equipment to avoid the risk of fire and burns.

Oxygen makes it easier for a fire to start and spread. Do not leave the nasal cannula or mask on bed coverings or chair cushions, if the device is turned on, but not in use; the oxygen will make the materials more flammable. Turn the device off when not in use to prevent oxygen enrichment.

It is the responsibility of the responsible organization to ensure that the oxygen source is compatible with the rated range of pressure, flowrate and oxygen concentration as marked on the equipment and indicated in the instructions for use as this can affect the performance of the equipment or pipeline system that can consequently result in serious deterioration of health.

Open flames during Optiflow high flow therapy are dangerous and are likely to result in fire or death. Do not allow open flames within 2 m of the equipment or any oxygen-carrying accessories.

Caution

Do not connect an oxygen supply to both the high-pressure oxygen inlet port and the low-pressure oxygen inlet port at the same time. Using the low-pressure inlet at the same time as the high-pressure inlet may cause incorrect oxygen delivery and a FiO_2 Above Target alarm.

! Note

The built-in oxygen analyzer uses ultrasonic measurement technology. It does not require in-field calibration.

4.4.1 High-pressure oxygen (HPO) source

When oxygen is connected to the HPO port, the Airvo 3 NIV directly controls the oxygen input to meet the target FiO_2 setting.

The High FiO_2 alarm can be disabled or a threshold between 30% and 95% can be selected when the Airvo 3 NIV is initially set up for your environment (see Oxygen high alarm threshold, Airvo 3 NIV Technical Manual). The alarm threshold is displayed on the Titrate FiO_2 screen, if enabled. Tap the FiO_2 tile to open the Titrate FiO_2 screen.

4.4.2 Low-pressure oxygen (LPO) inlet port

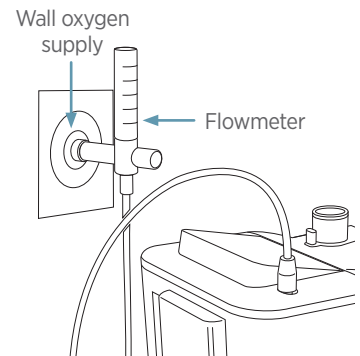
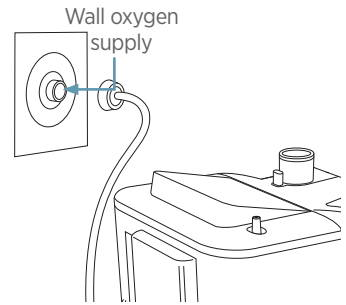
When using the LPO port, the amount of oxygen taken in by the Airvo 3 NIV is controlled by an external flowmeter.

Connect a tube from the external flowmeter to the LPO port. Make sure that the flowmeter is turned off whenever the Airvo 3 NIV is not delivering therapy.

When using the low-pressure oxygen inlet port, monitor the oxygen concentration displayed on the Home screen. The oxygen flow-regulator must be adjusted manually to maintain the prescribed oxygen concentration when changing the respiratory gas flow rate.

Clinicians may configure a High FiO_2 alarm to discourage use of high FiO_2 values in particular clinical environments.

The High FiO_2 alarm can be disabled or a threshold between 30% and 95% can be selected when the Airvo 3 NIV is initially set up for your environment (see Oxygen high alarm threshold, Airvo 3 NIV Technical Manual). The alarm threshold is displayed on the Titrate FiO_2 screen, if enabled. Tap the FiO_2 tile to open the Titrate FiO_2 screen.

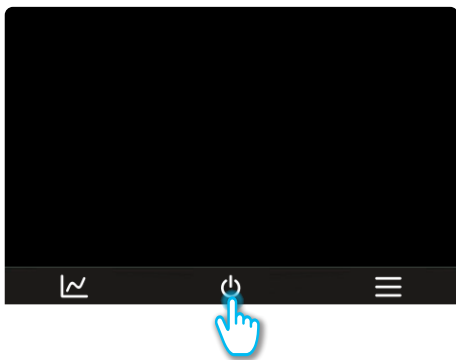


! Warning

Turn off the low-pressure oxygen source whenever the Airvo 3 NIV is not delivering therapy, to ensure that oxygen does not build up inside the device.

5. Using the Airvo 3 NIV

5.1 Getting started



Turn on the Airvo 3 NIV

Plug the Airvo 3 NIV power cord into the wall power supply.

Lock the wheels of the mobile pole stand to prevent the Airvo 3 NIV from moving.

Turn on the Airvo 3 NIV by holding down the Power on/off button for 2 seconds.

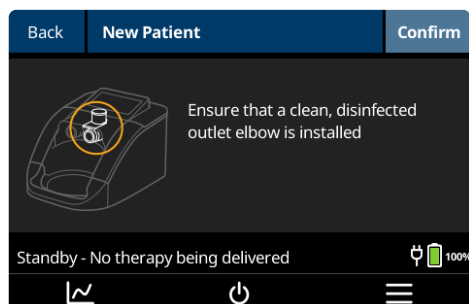
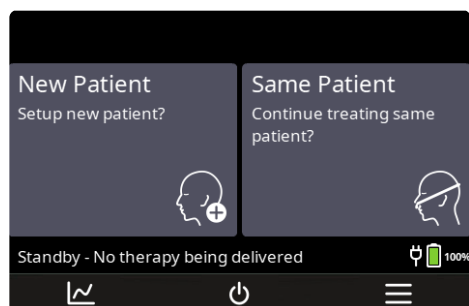
! Warnings

Make sure the Airvo 3 NIV is dry before plugging the power cord into the wall power supply to avoid a potential electric shock.

Do not cover or position next to materials that can block the gas intake, thereby interfering the patient therapy.

! Note

If the Airvo 3 NIV had been unused and disconnected from the wall power supply for some time, the device will not power on without being plugged in.

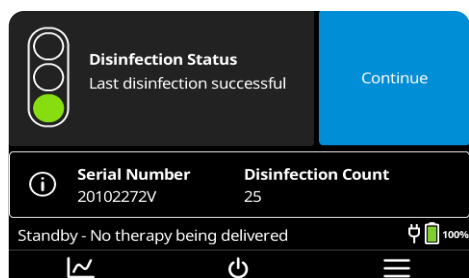


Review disinfection state

The Airvo 3 NIV will ask you if it will be used on:
the same patient who last used the device (tap Same Patient)
OR
a new patient (tap New Patient).

For a new patient, check that:

1. The outlet elbow has been cleaned and disinfected.
2. A new tube and chamber have been installed.



Review disinfection state (if the disinfection method is set to disinfection kit only)

For a new patient, check that:

1. The outlet elbow has been cleaned and disinfected.

The Airvo 3 NIV will indicate the outcome of the last disinfection cycle:



Green: The previous disinfection cycle was completed successfully.



Orange: A successful disinfection cycle has not been performed. Please run a successful disinfection cycle before use on a new patient.



Red: The previous disinfection cycle failed to complete. Please run a successful disinfection cycle before use on a patient.

The number of successful disinfection cycles completed by the Airvo 3 NIV is displayed in the lower left hand corner under 'Disinfection count'.

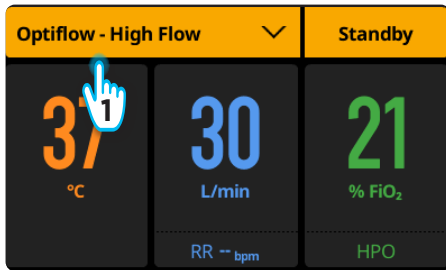
2. A new tube and chamber have been installed.

Warnings

The Airvo 3 NIV must be cleaned and disinfected between patients. Refer to section 8 for the steps required to reprocess the Airvo 3 NIV between patients.

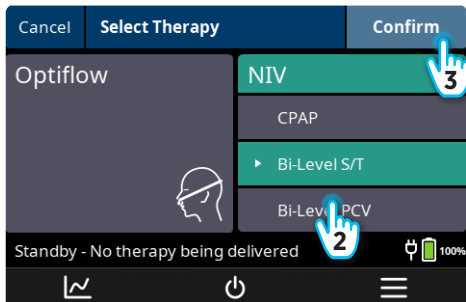
Do not exceed the maximum use period for single-patient-use accessories and consumables (see section 8.3 for the schedule for changing accessories).

5.2 Selecting the therapy



1. Select the required therapy if it is not shown in the therapy bar.
2. Select the prescribed therapy.
3. Tap Confirm to apply the change. Tap Cancel to discard any changes and return to the system menu.

If the Airvo 3 NIV is delivering therapy, tap the STOP button to enter standby mode. The therapy mode can not be changed while therapy is being delivered.



Note: Optiflow high flow and NIV require different tubes (see section 4.1). The Airvo 3 NIV will raise a Wrong Tube alarm if therapy is started with the wrong breathing tube connected.

5.3 Optiflow high flow therapy settings

The default range of Optiflow high flow therapy settings is shown below. Some settings may have been limited, or disabled, when the device was initially set up for its intended clinical environment. Refer to the Airvo 3 Technical Manual for details.

Settings are persistent and will retain their previous value when the Airvo 3 NIV is turned on. Selecting New Patient when reviewing the disinfection state (see section 5.1 above) applies the default values for its intended clinical environment to all settings.

Setting	Range	Description
Target humidity	31 – 37 °C	Target humidity for the respiratory gas supplied to the patient
Target flow	2 – 70 L/min	Flow rate of the respiratory gas supplied to the patient
FI _O ₂	21 – 100%	Target oxygen concentration for the breathing gases when an external oxygen supply is connected to the high-pressure oxygen inlet port
Expiratory relief (Target flow tile)	Off, 10%, 20%, 30%	This setting is disabled by default, and only available when the set flow is greater than 25 L/min, refer to the Airvo 3 NIV Technical Manual for details. Expiratory relief automatically reduces the respiratory gas flow rate during exhalation and returns it to normal during inhalation. Indicative flow rates are displayed on the settings screen. These may differ depending on the method and strength of the patient’s exhalation

Tiles on the Home screen show current Optiflow high flow therapy settings and measurements. Only tiles relevant to connected accessories are shown.

Therapy bar
Displays the active therapy. Tap to change the active therapy

Settings tiles
Displays therapy settings and measurements. Tap tiles to change settings or view more information

Messages and alarms
Displays therapy information and alarms

Optiflow - High Flow

Standby

37 °C

30 L/min

21 % FI_O₂

RR -- bpm

HPO

START

Standby - No therapy being delivered

Start therapy
Tap to start delivering therapy.

Status indicators

Data and Graphs

Power on/off

System menu

Target humidity setting

37 °C

Target flow setting (top)

Respiratory rate measurement † (bottom)

35 L/min

Oxygen concentration (FI_O₂) setting or measurement * †

21 % FI_O₂

SpO₂ ‡ (top)

Pulse rate (bottom)

SpO₂ alarm thresholds

93 % SpO₂

98 85

60 bpm

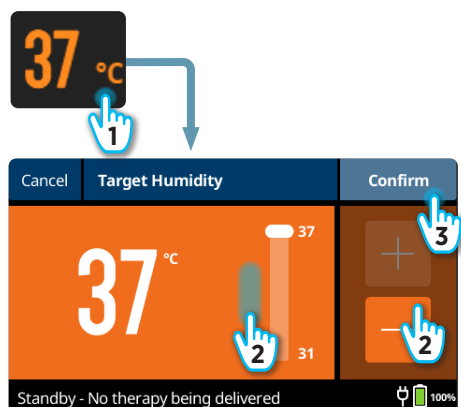
* The FI_O₂ tile shows the breathing gas oxygen concentration setting when supplementary oxygen is connected to the high-pressure oxygen (HPO) inlet port and measured oxygen concentration when connected to the low-pressure oxygen (LPO) inlet port. Measured oxygen concentration is not available in standby mode.

† "--" is displayed when a value is not available; values are gray when signal quality is poor.

‡ The SpO₂ tile is displayed automatically when a compatible pulse oximeter is connected.

5.4 Starting Optiflow high flow therapy

Follow the steps below to start delivering Optiflow high flow therapy. Some settings may have been limited, or disabled, when the device was initially set up for your clinical environment. Refer to the Airvo 3 NIV Technical Manual for details.



Adjust target humidity

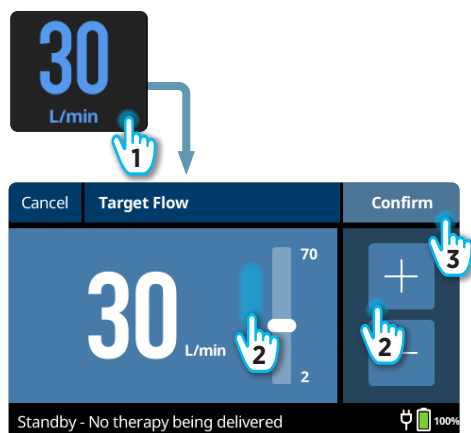
1. Tap the target humidity tile to open the Target Humidity screen.
2. Use the + / - buttons or slider to select a desired target humidity.
3. Tap Confirm to apply the change and return to the Home screen. Tap Cancel to discard any changes.

Warnings

Airvo 3 NIV is classified as a Category 1 humidifier for patients with bypassed airways (tracheostomies) in the following modes only: 37 °C and 10-60 L/min. Do not use any other mode for patients with bypassed airways (tracheostomies).

Note

Patients using face masks may find high temperatures uncomfortable. Consider a target temperature of 31 °C.



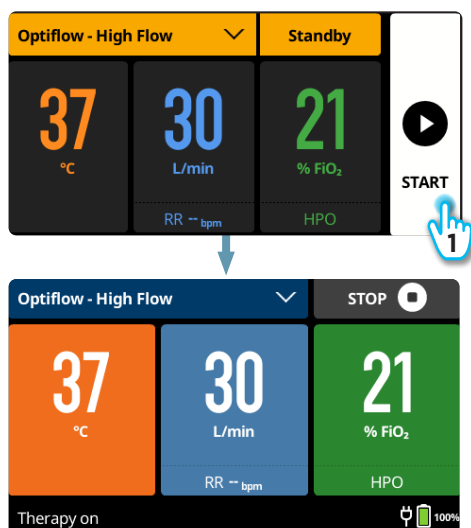
Adjust target flow

1. Tap the target flow tile to open the Target Flow screen.
2. Use the + / - buttons or slider to select the desired flow.
3. Tap Confirm to apply the change and return to the Home screen. Tap Cancel to discard any changes.

An appropriate flow rate for your patient should be prescribed following hospital protocols.

Note

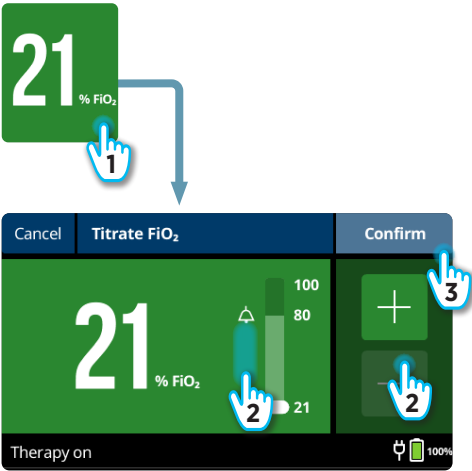
Refer to the patient interface user instructions for details.



Start therapy

Check that the breathing tube is assembled correctly and all connections are secure. Check the alarms are operating properly according to the instructions in section 7.5.

1. Tap the START button to begin therapy. After warming up, the Airvo 3 NIV will play a short melody and display the message "Therapy on".



Adjust supplementary oxygen (optional)

 Warnings

Use continuous SpO₂ monitoring on patients who would desaturate significantly if their oxygen supply is disrupted.

The FiO₂ control limits should be prescribed based on patient condition, hospital policies and clinical judgement for Optiflow nasal high flow therapy.

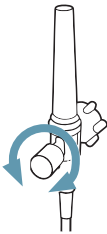
Oxygen connected to the high-pressure inlet port (HPO)

1. Tap the FiO₂ tile to open the Titrate FiO₂ screen.
2. Use the + / - buttons or slider to select the desired FiO₂.
3. Tap Confirm to apply the change and return to the Home screen. Tap Cancel to discard any changes.

The Airvo 3 NIV will automatically adjust oxygen flow to maintain the selected FiO₂.

Oxygen connected to the low-pressure inlet port (LPO)

The Airvo 3 NIV does not directly control FiO₂.
Use an external flowmeter to adjust FiO₂ to the prescribed level.
The oxygen tile displays measured FiO₂.



 Note

It may take a few minutes for the oxygen measurement to stabilize.

The external flowmeter will need to be readjusted following changes to Airvo 3 NIV target flow.

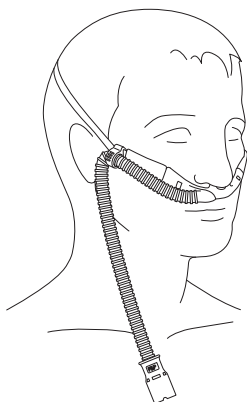
High FiO₂ alarm

Clinicians may configure a High FiO₂ alarm to discourage the use of high FiO₂ values in particular clinical environments. See the Airvo 3 Technical Manual for setup details.

If the alarm is enabled, the alarm threshold is displayed on the Target FiO₂ screen.

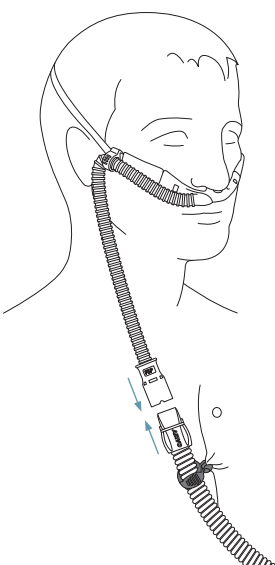
The following table shows the target flow settings able to be used with these interfaces

		L/min																	
PATIENT INTERFACE		2	3	4	5	6	7	8	9	10	15	20	25	...	50	55	60	65	70
Optiflow Junior 2	OJR414 M	2				7													
	OJR416 L	2										20							
	OJR418 XL	2											25						
	OJR520 XXL										10				50				
Optiflow+ Interfaces	OPT942/OPT962 /OPT1042 (S)										10						60		
	OPT944/OPT964/ OPT1044 (M)										10							70	
	OPT946/OPT966/OPT1046 (L)										10								70
												10							
Trache	OPT970										10						60		
Mask Adapter	OPT980										10						60		



Fit the patient interface

Fit the patient interface to your patient following the user instructions supplied with the interface. Take care to follow all warnings and cautions.



Connect to the patient interface

Connect the patient interface to the connector at the end of the breathing tube.

The patient may be connected to the heated breathing tube immediately. When therapy initiation is not urgent, it is recommended to wait until the Airvo 3 NIV plays a short melody and displays "Therapy On" in the Message bar.

Attach the breathing tube clip to the patient's clothing.

Warnings

Do not allow the breathing tube to remain in direct contact with the patient's skin for long periods of time to avoid the risk of burns. The healthcare professional shall assess the conditions for safe contact, such as duration and skin condition.

Do not cover or add heat above ambient levels to any part of the breathing tube or interface e.g. by covering with a blanket or by heating with infrared radiation, an overhead heater, or an incubator, as this can affect the quality of the therapy or injure the patient



Do not use sealed patient interfaces with Optiflow high flow therapy, to avoid the risk of suffocation or barotrauma.

Ensure a sufficient intended leakage between the breathing system and the patient to allow the patient to exhale.

Caution

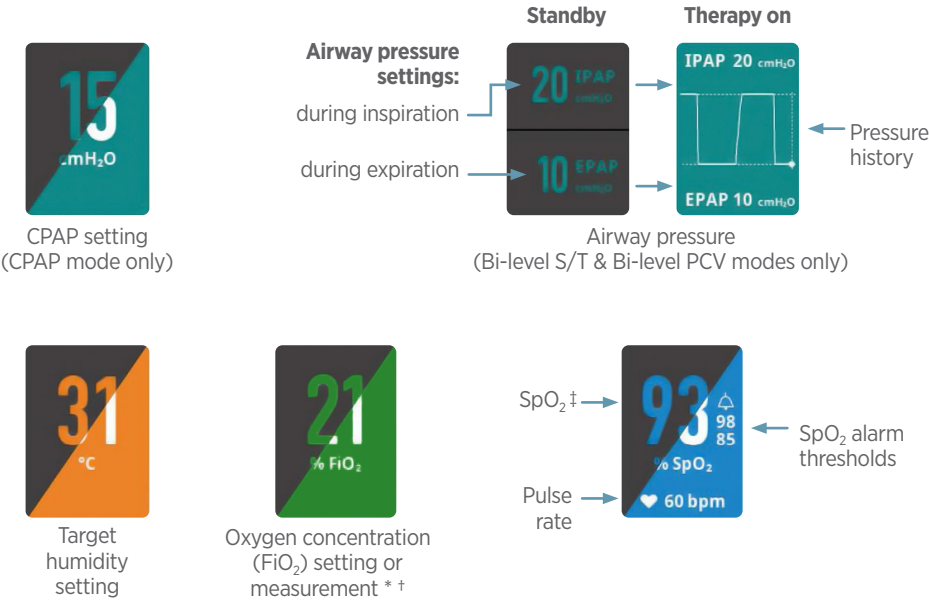
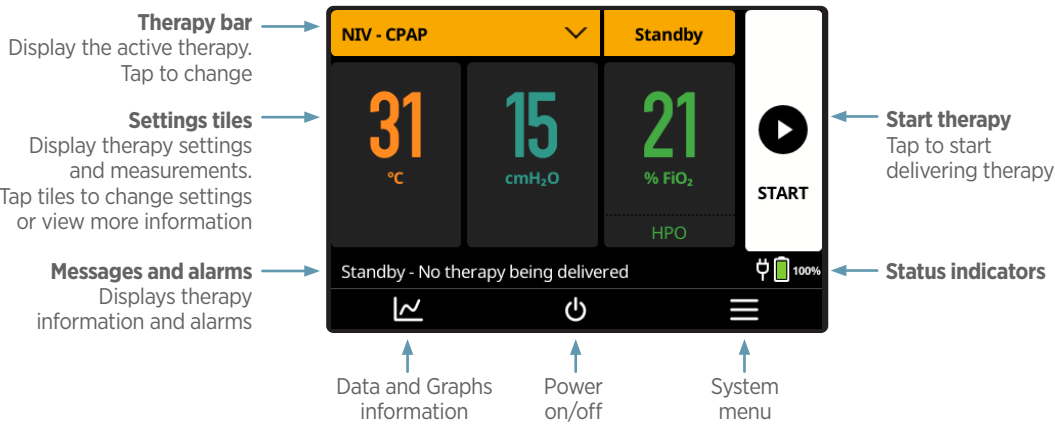
Keep the heated breathing tube away from electrical monitoring leads (e.g. EEG, ECG/EKG, EMG, pulse oximeter) to reduce the risk of interference with the signal monitored.

Note

The air may feel warm when your patient starts using the Airvo 3 NIV. This is normal. The patient should continue to breathe normally.

5.5 Adult NIV therapy settings

Tiles on the Home screen show current NIV therapy settings and measurements. Only tiles relevant to connected accessories and the selected therapy mode are shown.



* The FiO₂ tile shows the breathing gas oxygen concentration setting when supplementary oxygen is connected to the high-pressure oxygen (HPO) inlet port and measured oxygen concentration when connected to the low-pressure oxygen (LPO) inlet port. Measured oxygen concentration is not available in standby mode.

† "--" is displayed when a value is not available; values are gray when signal quality is poor.

‡ The SpO₂ tile is displayed automatically when a compatible pulse oximeter is connected.

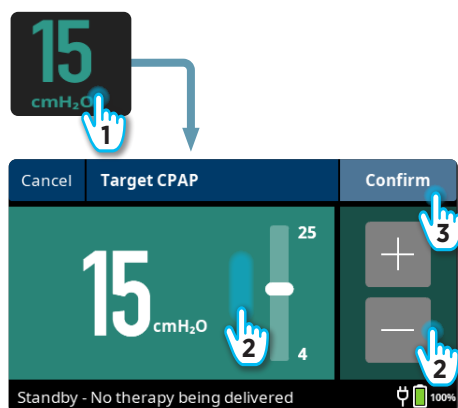
5.6 Starting adult NIV therapy

Follow the steps below to start delivering NIV therapy. Some settings may have been limited, or disabled, when the device was initially set up for your clinical environment. Refer to the Airvo 3 NIV Technical Manual for details.



Adjust target humidity

1. Tap the Target humidity tile to open the Target Humidity screen.
2. Use the + / - buttons or slider to select a desired target humidity.
3. Tap Confirm to apply the change and return to the Home screen. Tap Cancel to discard any changes.



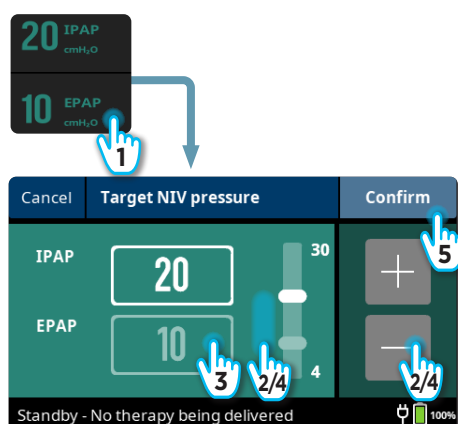
Adjust airway pressure

CPAP mode

Change the airway pressure to match the CPAP value prescribed for your patient:

1. Tap the CPAP tile to open the Set CPAP screen.
2. Use the + / - buttons or slider to change the CPAP value.
3. Tap Confirm to apply the change and return to the Home screen. Tap Cancel to discard any changes.

You can select a constant airway pressure between 4 cmH₂O and 25 cmH₂O.



Bi-level S/T and Bi-level PCV modes

Change the inspiratory and expiratory airway pressures to match the values prescribed for your patient:

1. Tap the IPAP tile to open the Set NIV Pressure screen.
2. Use the + / - buttons or slider to change the IPAP value.
3. Tap the EPAP value to select it.
4. Use the + / - buttons or slider to change the EPAP value.
5. Tap Confirm to apply the change and return to the Home screen. Tap Cancel to discard any changes.

You can select EPAP between 4 cmH₂O and 25 cmH₂O and IPAP between 4 cmH₂O and 30 cmH₂O. EPAP cannot be set higher than IPAP.

The settings for NIV therapy are shown below. Some settings may have been limited, or disabled, when the device was initially set up for your clinical environment. Refer to the Airvo 3 NIV Technical Manual for details.

The settings for NIV therapy are shown below. Some settings may have been limited, or disabled, when the device was initially set up for your clinical environment. Refer to the Airvo 3 NIV Technical Manual for details.

Name	Menu†	Mode‡	Range	Description
Target Temperature		C B P	25 – 31 °C	Target temperature for the respiratory gas supplied to the patient.
CPAP		C	4 – 25 cmH ₂ O	Positive airway pressure for the complete breathing cycle.§
IPAP		B P	4 – 30 cmH ₂ O	Positive airway pressure during inspiration.*§
EPAP		B P	4 – 25 cmH ₂ O	Positive airway pressure during expiration/rest.*§
FiO₂		C B P	21 – 100%	Oxygen concentration of the breathing gases when supplementary oxygen is connected to the high-pressure oxygen inlet port.
Ramp		C B P		Access the ramp settings (see below).
Backup Rate		B P	4 – 60 breaths/min	Rate of machine triggered breaths imposed when patient breaths are not detected. The backup rate is limited to a maximum of 0.5 * (60/Inspiration time).
Inspiration Time		B	0.3 – 4.0 s	The duration of IPAP phase for machine-triggered breaths. The inspiration time is limited to a maximum of 0.5 * (60/Backup rate).
Inspiration Time		P	0.3 – 4.0 s	The duration of IPAP phase for machine-triggered breaths, and the imposed duration of patient-triggered breaths. The inspiration time is limited to a maximum of 0.5 * (60/Backup rate).
Rise Time		B P	0.2, 0.3, 0.5, 0.7 or 0.9 s	The time for pressure to rise from EPAP to IPAP when a breath is triggered. Adjust to find the most comfortable setting for your patient.
Triggering		B P	2 – 9 L/min	Sensitivity to patient inhalation for detecting breaths. Lower values are more sensitive.
Mask type		C B P	F&P NIV / F&P OptiNIV / Other	The type of patient interface mask used to deliver therapy.

† The setting location in the Airvo 3 NIV user interface is:



Home screen



NIV Advanced Settings menu. Tap , then select NIV Advanced Settings

‡ Modes are abbreviated as C: CPAP, B: Bi-level S/T, P: Bi-level PCV, and indicate the therapy modes where each setting applies.

* The EPAP value cannot be set higher than the IPAP value. The IPAP value cannot be set lower than the EPAP value.

§ Pressures are measured above atmospheric pressure (gauge pressure). The Airvo 3 NIV does not generate sub-atmospheric pressures.

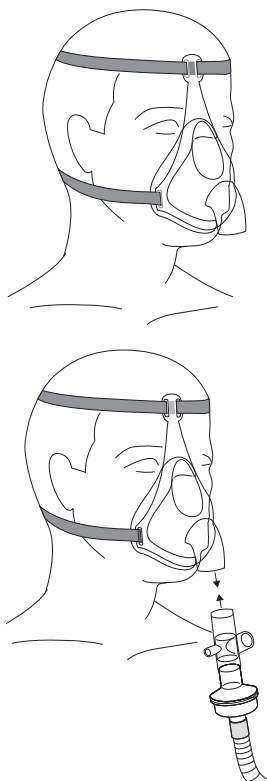
Select Ramp from the NIV Advanced Settings menu to configure the pressure ramp applied when therapy begins.

Name	Mode†	Range	Description
Enable Ramp	C B P	On/Off	Enable/disable ramp mode. The Ramp button will be displayed on the Home screen when ramp mode is enabled.
Ramp Time	C B P	5 – 45 min	Duration of the ramp from the initial airway pressure target to the final pressure target (when the pressure ramp is on).
Start CPAP	C	4 – 25 cmH ₂ O	CPAP value at the start of a ramp (when the pressure ramp is on).§
Start IPAP*	B P	4 – 30 cmH ₂ O	IPAP value at the start of a ramp (when the pressure ramp is on).§
Start EPAP*	B P	4 – 25 cmH ₂ O	EPAP value at the start of a ramp (when the pressure ramp is on).§

† Modes are abbreviated as C: CPAP, B: Bi-level S/T, P: Bi-level PCV, and indicate the therapy modes where each setting applies.

§ Pressures are measured above atmospheric pressure (gauge pressure). The Airvo 3 NIV does not generate sub-atmospheric pressures.

* A Start IPAP/EPAP setting cannot be set higher than the final IPAP/EPAP setting.



Connect the patient interface and connect the breathing tube

Fit the patient interface to your patient following the user instructions supplied with the interface. Take care to follow all warnings and cautions.

Connect the patient-interface connector to the connector at the end of the breathing tube.

Warnings

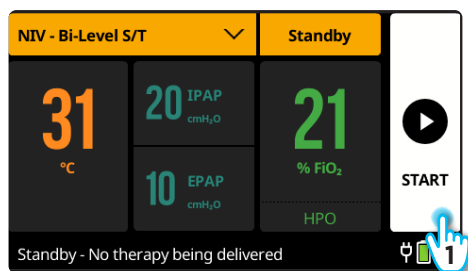
If the breathing tube filter is missing, ruptured or faulty the entire gas pathway may become contaminated with exhaled respiratory gas and/or bodily fluids under extreme conditions.

Use only compatible Fisher & Paykel Healthcare breathing tube filters. Incompatible filters could affect the quality of therapy or injure the patient.

Do not modify the breathing tube filter or exhalation port in any way. Modifications may cause patient injury or reduce the quality of therapy.

Caution

Keep the heated breathing tube away from electrical monitoring leads (e.g. EEG, ECG/EKG, EMG, pulse oximeter) to reduce the risk of interference with the signal monitored.



Start therapy

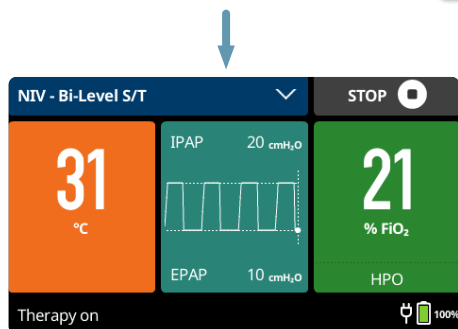
Check that the breathing tube is assembled correctly and all connections are secure. Check the alarms are operating properly according to the instructions in section 7.5.

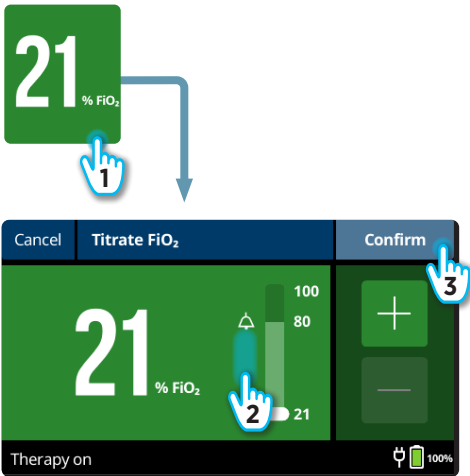
1. Tap the START button to begin therapy. After warming up, the Airvo 3 NIV will play a short melody and display the message "Therapy on".

Warnings

Check that the NIV alarm settings are appropriate for the patient.

Use continuous SpO₂ monitoring on patients who would desaturate significantly if their oxygen supply is disrupted.





Adjust supplementary oxygen (optional)

Oxygen connected to the high-pressure inlet port (HPO)

1. Tap the FiO₂ tile to open the Titrate FiO₂ screen.
2. Use the + / - buttons or slider to adjust the oxygen concentration between 21 and 100%.
3. Tap Confirm to apply the change and return to the Home screen. Tap Cancel to discard any changes.

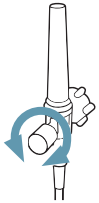
The Airvo 3 will automatically adjust oxygen flow to maintain the selected FiO₂.

Oxygen connected to the low-pressure inlet port (LPO)

The Airvo 3 NIV does not directly control FiO₂. Use an external flowmeter to adjust FiO₂ to the prescribed level. The Oxygen tile displays measured FiO₂.

Note

It may take a few minutes for the oxygen measurement to stabilize. The external flowmeter will need to be readjusted following changes to the Airvo 3 NIV target flow.

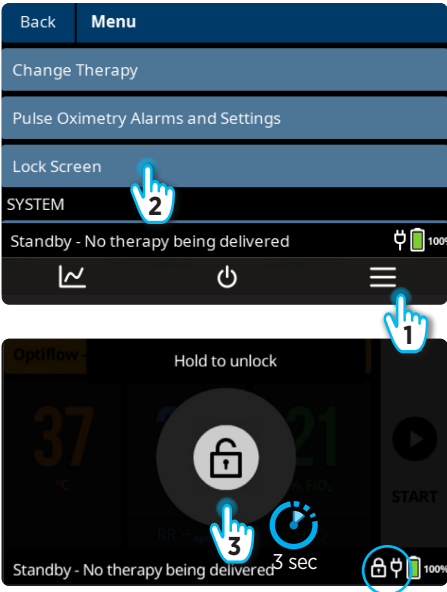


5.7 During therapy

Monitor the patient following hospital protocols and clinical judgement. Ensure you can hear and respond to any device alarms. If there is an interruption to the power supply, and the battery is depleted, the Airvo 3 NIV will raise a Power Out alarm, turn off, and not deliver any therapy to the patient. The power out alarm will sound once every 10 seconds for a minimum of 120 seconds, and the signal light above the touch screen will flash. Once power is restored the Airvo 3 NIV can be restarted and will retain the previous therapy and alarm settings

5.7.1 Lock screen (optional)

The lock screen can prevent accidental settings changes.



To enable the touch lock:

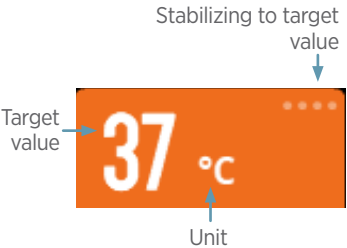
1. Tap ≡ to open the system menu.
 2. Select Lock screen from the system menu.
- The symbol  is shown in the Message bar.

To disable the touch lock:

3. Touch the screen while it is locked, then hold the Unlock icon for three seconds.

5.7.2 Monitor and adjust settings

Adjust settings as needed. Most changes take effect after pressing the confirmation button but it may take a few minutes for some settings, such as target humidity, to respond to changes. Tiles show an animated ellipsis symbol (...) to indicate that a therapy setting has not yet reached its target.



5.7.3 Manage condensation

Drain excess condensate from the breathing tube by:

1. Disconnecting the breathing tube from the patient interface (stop therapy first if in NIV mode), and
2. Lifting the patient end of the tube so the condensate runs into the water chamber.

If in high flow mode, reduce the flow rate below 30 L/min if the condensate does not run freely into the water chamber. Return the flow rate to the prescribed setting after draining the breathing tube.

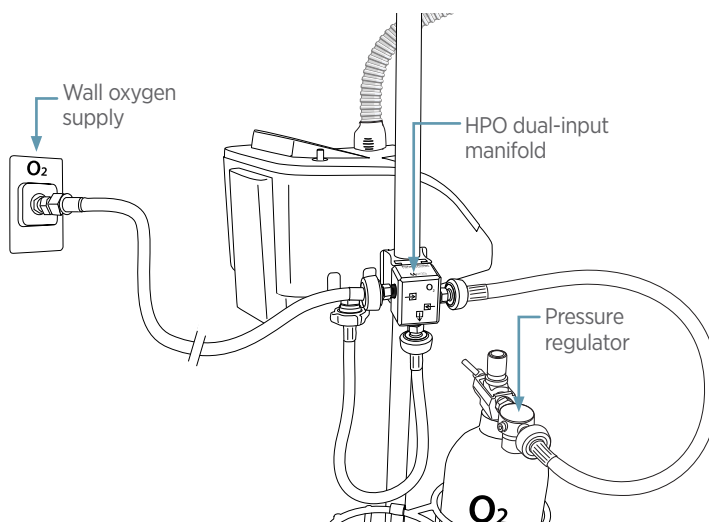
Direct cold air away from the heated breathing tube where possible. Air conditioners, fans, open windows and other sources of cold air may increase condensation. Consider reducing the target humidity if condensation persists.

5.8 Mobility and battery operation

The HPO (High Pressure Oxygen) dual-input manifold and internal rechargeable battery provide continuity during intra-hospital transport. Reduced humidity will be delivered when the Airvo 3 NIV is being powered only by the battery; for more details see Appendix 4. HPO dual-input manifold uses the oxygen supply with the highest pressure.

When transporting the Airvo 3 NIV with your patient:

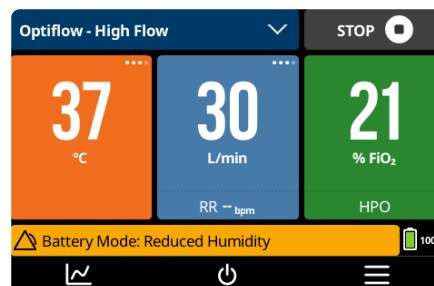
1. Adjust therapy settings as necessary for intra-hospital transport.
2. If using supplementary oxygen:
 - Check that the oxygen bottle contains enough oxygen for your journey.
 - Turn on the oxygen-bottle pressure regulator.
 - Disconnect the oxygen hose from the wall supply. Either attach it to a second oxygen bottle for longer trips or hook it over the Airvo mobile pole stand if additional oxygen is not required.



The HPO dual-input manifold will use the oxygen-bottle supply automatically. Check the battery contains enough charge for intra-hospital transport. A new battery will provide therapy for approximately 40 minutes when fully charged. A Low Battery alarm will occur when 35% of the battery is remaining (no changes to the device or therapy). A Critically Low Battery alarm will occur when 20% of the battery is remaining (humidity is turned off, oxygen and flow continue to be delivered). When the battery is fully depleted, the Airvo 3 NIV will interrupt therapy and produce a Power Out alarm.

3. Unplug the Airvo 3 NIV from the wall power.
4. The Airvo 3 NIV will display a Battery Mode: Low Humidity alarm.
5. When you reach your destination:
 - Reconnect the Airvo 3 NIV to the wall power supply and the wall oxygen supply.
 - Turn off the oxygen bottle pressure regulator to avoid draining the oxygen bottle and switch to the wall oxygen supply.

If you are not using the oxygen dual-input block, connect an oxygen bottle (if required) to one of the oxygen inlet ports when transporting your patient. Ensure any oxygen supply connected to the low-pressure oxygen (LPO) inlet port is turned off when the device is in standby mode, not delivering therapy.



Warning

If using the battery as the power source, periodically check the battery status to ensure the battery does not become depleted while therapy is being delivered.

Warnings

Only use the Airvo 3 NIV battery with the Airvo 3 NIV device.

Only charge the Airvo 3 NIV battery with the Airvo 3 NIV device.

Loss of power will result in loss of therapy. In the event of a Critically Low Battery alarm, promptly connect the Airvo 3 NIV to the wall power supply to avoid loss of therapy due to the battery becoming depleted.

Contact technical personnel to remove the battery from the device if it is not likely to be used for an extended period of time.

5.9 Changing therapy

The Airvo 3 NIV supports both Optiflow high flow and noninvasive ventilation therapies in a single device. Switching the therapy provided to a patient involves changing the breathing tube and patient interface, then selecting the new therapy on the device. When alternating between therapies, store unused patient interfaces and heated breathing tubes in a clean environment for future use on the same patient.



1. Stop therapy

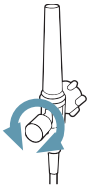
If therapy is being delivered, stop therapy by tapping the STOP button.

Note

The Airvo 3 NIV will raise a Tube Missing alarm if you disconnect the breathing tube before stopping therapy. Tap the Audio Pause button, or reconnect the breathing tube and tap the Stop button.

2. Remove the patient interface

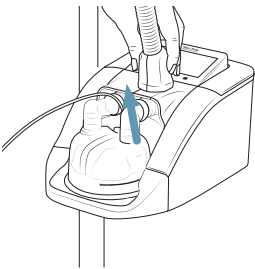
Remove the interface from your patient and disconnect it from the breathing tube.



3. Turn off supplementary oxygen

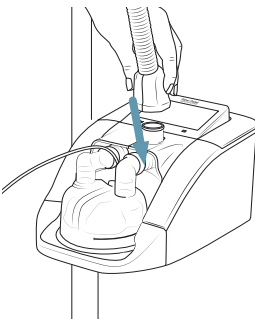
Turn off supplementary oxygen using the flow regulator if oxygen is added through the low-pressure oxygen inlet port on top of the Airvo 3 NIV.

It is not necessary to disconnect oxygen from the high-pressure oxygen inlet port.



4. Disconnect the heated breathing tube

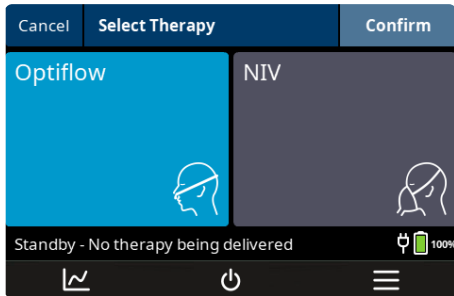
Disconnect the heated breathing tube from the Airvo 3 NIV by squeezing the sides of the connector and pulling up.



5. Connect the heated breathing tube

Connect the appropriate heated breathing tube for the new therapy:

- For NIV therapy, assemble the NIV breathing tube following the instructions in section 4.3. Be sure to fit the filter, and exhalation port correctly (if required).
- For Optiflow high flow therapy, assemble the heated breathing tube following the instructions provided with the interface.



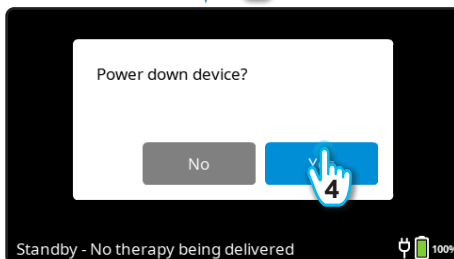
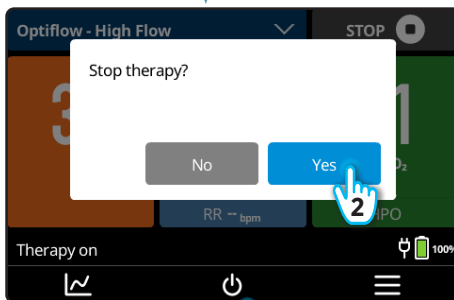
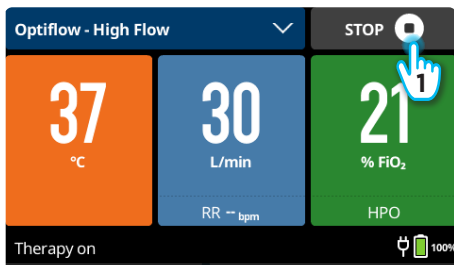
6. Select the new therapy

Follow the instructions in section 5.2 to select the new therapy.

Then continue with:

- section 5.4 for Optiflow High Flow therapy, or
- section 5.5 for adult NIV therapy.

5.10 Stopping therapy



When therapy is finished:

1. Remove the patient interface from your patient.
2. If oxygen is provided through the low-pressure oxygen inlet port on the top of the Airvo 3 NIV, turn off and disconnect the oxygen supply.

ⓘ Note

The Airvo 3 NIV will automatically stop oxygen provided through the high-pressure oxygen inlet port. You do not need to disconnect it.

1. Tap the STOP button to end therapy.
2. Review any warnings, then tap Yes to confirm and enter standby mode or No to continue therapy.
3. Turn the Airvo 3 NIV off by holding down the Power button for 2 seconds.
4. Tap Yes to power down the device.

The Airvo 3 NIV must be cleaned and disinfected between patients. Follow the reprocessing instructions if your patient has finished using the device.

⚠ Warnings

To avoid burns, do not touch the heater-plate or the bottom of the water chamber. The water in the chamber and the heater-plate beneath the chamber become hot during use.

Turn off the low-pressure oxygen source before stopping therapy. The oxygen flow must be turned off when the Airvo 3 NIV is not delivering therapy to ensure oxygen does not build up inside the device.

6. Monitoring data

 Warning

In line with the indications for use of the Airvo 3 NIV, the monitoring functionality of the Airvo 3 NIV is intended for use on spontaneously breathing patients and not intended for patients requiring life support. It is the responsibility of the clinician to select the appropriate level of monitoring for their patient and to be prepared to deal with alarms and equipment malfunction. Additional, independent monitoring equipment may be necessary.

The Airvo 3 NIV records up to 24 hours of data for review on the Data and Graphs screen, accessible by tapping the Data and Graphs Information button  from the Home screen. Data and Graphs data will be lost if power from the battery and from the wall supply is lost. Refer to the Airvo 3 NIV Technical Manual for detailed information on data handling.

The Airvo 3 NIV is designed not to collect identifiable information about end-users. To function effectively, Airvo 3 NIV will collect and store limited therapy data.

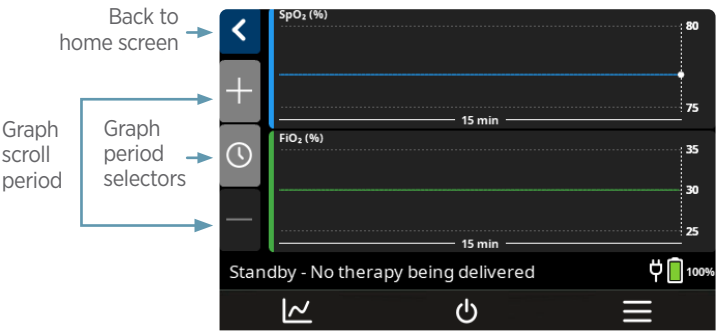
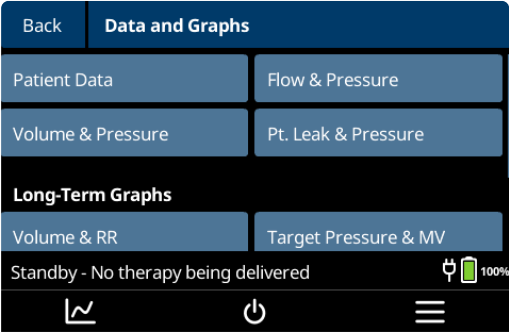
Limited Airvo 3 NIV device information may be collected by F&P to monitor medical device performance, including device identifiers. This is to monitor medical device effectiveness, and improvement opportunities (e.g. firmware). Information is stored and used securely by F&P and does not include any data relating to your patient's personal information.

For more information about what type data is involved in these activities, refer to the Airvo 3 Technical Manual.

Please refer to the T&Cs for your data protection and privacy obligations. Alternatively, refer our Global Privacy Statement on our website for more on how we handle personal information.

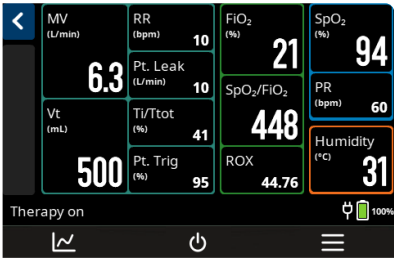
Data can be transferred from the Airvo 3 NIV to data collection devices via the data port adapter. Integration with the data collection device is required.

The graph selector screen shows all of the available graphs for the therapy. A graph can be accessed by pressing on a tile. Graphs can alternatively be selected using the scroll button outside of the graph selector screen.



6.1 Patient data

The values displayed in the Patient data screen are described below. Measurements that are not available are shown as “-”. Measurements may not be available when the Airvo 3 NIV is in standby mode or the device has not collected enough data for a reliable measurement.



Therapy	Label	Unit	Description
Optiflow high flow	Flow	L/min	The current flow rate of breathing gases supplied to the patient
Optiflow high flow	RR	BPM	The patient's respiratory rate (breaths per minute), averaged over the last 90 seconds
NIV	Patient leak	L/min	The estimated unintentional leak rate (BTPS) for the last breath. Displayed when Mask Type is 'F&P' or 'OptiNIV'
NIV	Total leak	L/min	The estimated combined intentional and unintentional leak rate (BTPS). Displayed when Mask Type is 'Other'
NIV	MV	L/min	Effective minute ventilation (BTPS) estimated from a moving average over the last 5 breaths
NIV	RR	BPM	The respiratory rate (breaths per minute), averaged over the last 5 breaths. Includes both machine and patient triggered breaths
NIV	Pt. Trig	%	The proportion of breaths triggered by the patient relative to the total number of breaths supplied over the last 15 minutes. Additional breaths are triggered at the Backup rate
NIV	Ti/Ttot	%	Inspiration duty cycle: the proportion of time spent at IPAP relative to the total breathing cycle displayed as a moving average for the last 5 breaths. (Not available in CPAP mode)
NIV	Vt	mL	Estimated expiratory tidal volume, displayed as a moving average over the last 5 breaths. Compensated for leak volume, body temperature, pressure and respiratory humidity (BTPS)
Optiflow high flow NIV	Temp	°C	The current humidity of breathing gases supplied to the patient
Optiflow high flow NIV	FiO ₂	%	The current fraction of oxygen in breathing gases supplied to the patient
Optiflow high flow NIV	SpO ₂ /FiO ₂		Ratio of SpO ₂ and FiO ₂
Optiflow high flow NIV	ROX		Ratio of SpO ₂ /FiO ₂ to respiratory rate
Optiflow high flow NIV	SpO ₂	%	Peripheral blood oxygen saturation measured by pulse oximeter
Optiflow high flow NIV	PR	BPM	Pulse rate measured by pulse oximeter (beats per minute)

6.2 Live value graphs

Airvo 3 NIV live value trends graphs show the last 30 seconds of data.

Therapy	Label	Unit	Description
Optiflow high flow NIV	Flow	L/min	The current flow rate of breathing gases supplied to the patient
NIV	Pressure	cmH ₂ O	Airway pressure
NIV	Tidal volume	mL	Estimated expiratory tidal volume. Compensated for leak volume, body temperature, pressure and respiratory humidity (BTPS)
NIV	Patient Leak (Pt.Leak)	L/min	Estimated unintentional leak rate (BTPS). Displayed when Mask Type is 'F&P' or 'OptiNIV'
NIV	Total Leak (TTL. Leak)	L/min	Estimated total leak (BTPS), it is the combined intentional and unintentional leak rate. Displayed when Mask Type is 'Other'

6.3 Long term graphs

Airvo 3 Data and Graphs show measurements plotted against time for up to 24 hours. New measurements are added to the right side of the graph. Prior data will scroll to the left as new measurements are added. Gaps will appear in the plotted data if therapy is stopped or measurements are missing due to poor signal quality.

The graphs available are described in the table below.

Therapy	Label	Unit	Description
Optiflow high flow	Flow	L/min	The set flow rate of breathing gases supplied to the patient
Optiflow high flow	RR	BPM	The patient's respiratory rate (breaths/minute), averaged over the last 90 seconds
NIV	MV	L/min	Effective minute ventilation estimated from a moving average over the last 5 breaths
NIV	RR	BPM	The patient's respiratory rate (breaths per minute), averaged over the last 5 breaths
NIV	Vt	mL	Estimated expiratory tidal volume, displayed as a moving average over the last 5 breaths. Compensated for leak volume, body temperature, pressure and respiratory humidity (BTPS)
NIV	Pressure	cmH ₂ O	Airway pressure settings
Optiflow high flow NIV	FiO ₂	%	The fraction of oxygen in breathing gases supplied to the patient
Optiflow high flow NIV	SpO ₂ /FiO ₂		Ratio of SpO ₂ and FiO ₂
Optiflow high flow NIV	ROX		SpO ₂ divided by FiO ₂ and respiratory rate
Optiflow high flow NIV	SpO ₂	%	Peripheral blood oxygen saturation measured by pulse oximeter
Optiflow high flow NIV	PR	BPM	Pulse rate measured by pulse oximeter (beats per minute)

7. Troubleshooting

This section describes common causes, and solutions to, problems and alarms that you may encounter while using the Airvo 3 NIV. The Airvo 3 Technical Manual contains additional information that may be helpful in resolving more advanced problems.

7.1 Alarms

The Airvo 3 NIV has visual and auditory alarms to notify users about interruptions to a patient's treatment. These alarms are generated by an intelligent alarm system, which processes information from sensors and target settings of the device and compares this information with pre-programmed limits. Changes to alarm settings will be preserved during or after any power loss.

The signal light flashes and troubleshooting information is displayed on the Airvo 3 NIV touch screen when any alarm is active. The color of the signal light indicates the highest-priority active alarm condition.



7.2 Alarm priority

Alarms are grouped by urgency and severity into three priority levels: low, medium, high. When multiple alarms are active, the audible alert, signal light and Message bar background color will signal the highest-priority alarm active.

- A response is needed for all alarms.
- A prompt response is required for all medium-priority alarms.
- An immediate response is required for all high-priority alarms.

Priority	Message bar, signal light color	Audible alert
Low	Solid yellow	High then low-pitched beep
Medium	Flashing yellow	3 beeps every 9 seconds
High	Flashing red	10 beeps every 5 seconds



Warning

Audible alarms may not be heard if the alarm volume is set lower than ambient noise. Missed alarms may lead to patient injury. Refer to the Airvo 3 Technical Manual to review and set the alarm volume.

7.3 Auditory information signals

The informative sounds made by the Airvo 3 NIV are:

Melody	Meaning
Ascending sequence of 5 tones	The breathing gas has warmed up
Single tone	A touch on the display has been registered
Single low then high tone	All active alarms have been resolved
High note followed by 2 (identical) lower notes, repeated every 10 seconds	The power out alarm is active. The wall power supply has been disconnected or turned off and the (optional) battery is depleted
Descending sequence of 3 tones	The device has completed the power off process
Sequence of 3 tones with high, low then middle pitch	The device has been turned on

7.4 Viewing alarm details

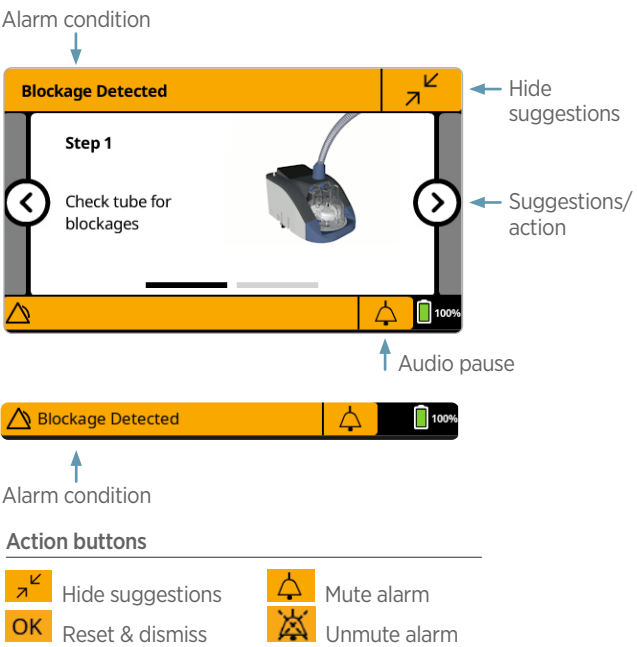
Alarms are displayed with suggestions and action buttons for managing the alarm information:

- Tap the Audio Pause button to silence the alarm for 120 seconds. The Audio Pause button will change to  when audible alarms are silenced.
- Use   to scroll through multiple suggestions. Some alarms have only one suggested resolution.
- Tap Hide suggestions to collapse the alarm information to the Message bar. Restore suggestions by tapping the alarm condition on the Message bar.

The alarm condition and action button are displayed on the Message bar when alarm information is collapsed.

The Message bar cycles through each alarm condition when multiple alarms are active. Tapping the Message bar displays a list of active alarm conditions, from highest priority to lowest priority and they are ordered from when they occurred.

Alarm signals always indicate the highest-priority active alarm condition.

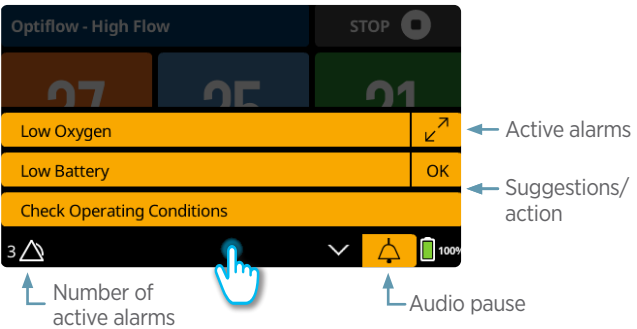


7.5 Checking the alarm system

To test the alarm system:

1. In standby mode, disconnect the breathing tube then press “Start”.
2. Verify that the “Check tube” visual alarm appears on screen.
3. Verify that the signal light flashes yellow.
4. Verify that the auditory alarm signal can be heard.

Do not use the Airvo 3 NIV if it fails this test. Contact your Fisher & Paykel Healthcare representative.



7.6 Airvo 3 NIV alarms

All the alarms you may encounter while using the Airvo 3 NIV are listed below with common causes, resolutions and any delays inherent in determining alarm conditions. These priorities have been allocated for an intended operator’s position of 2 meters from the device. The Airvo 3 NIV also uses an internal-priority ranking system. If multiple alarm conditions occur simultaneously, the device will display the highest-priority alarm.

Alarm condition	Priority	Delay	Meaning
Faults			
Device Fault [Fault X.X.X]	High	-	A technical fault has occurred that has caused therapy to be stopped. Restart the device to resolve the fault. If the issue persists, contact your service representative.
Device Fault [Fault X.X.X]	Med.	-	A technical fault has occurred, and the device is able to continue supplying limited therapy. Restart the device to resolve the fault. If the issue persists, contact your service representative.
Power alarms			
Power Out	High	< 5 s	The Airvo 3 NIV has been disconnected from the wall power supply and the internal battery is depleted. The auditory alarm will sound once every 10 seconds for a minimum of 120 seconds and the signal light above the touch screen will flash. The touch screen is off during the Power Out alarm. The Airvo 3 NIV will shut down after signaling the Power Out alarm but will restart automatically if power is restored before it shuts down.

Alarm condition	Priority	Delay	Meaning
Unsupported Battery	Med.	< 5 s	The device is running off the battery and either an incorrect battery type is connected or communications with the battery could not be established. Charging is disabled. During battery use the behavior is the same as the Critically Low Battery alarm
Critically Low Battery	Med.	< 5 s	The Airvo 3 NIV battery level is critically low and indicates at least 5 minutes left for complete loss of battery power. The humidification is turned-off to maintain operation of the blower and oxygen supply. Connect to wall power supply to continue therapy as normal.
Low Battery	Low	< 5 s	The Airvo 3 NIV battery level is low and indicates at least 10 minutes left for complete loss of battery power. Connect to wall power supply to continue therapy as normal.
Battery Mode: Reduced Humidity	Low	< 5 s	The Airvo 3 NIV has been disconnected from the wall power supply and the device is now running off the battery. The delivered humidity may be reduced.
Battery Charger Fault	Low	< 30 s	The battery charger is not functioning correctly and has been disabled. Restart the device to resolve the fault. If the issue persists, contact your service representative.
Therapy alarm - tube			
Outlet Elbow Missing	High	< 15 s	The Airvo 3 NIV outlet elbow has been removed from the device during therapy. Check that the outlet elbow is fully inserted into the Airvo 3 NIV. If the issue persists, replace the outlet elbow.
Check Tube	Med.	< 5 s	The Airvo 3 NIV cannot detect the heated breathing tube. Check that the heated breathing tube is not damaged and is plugged in correctly. Replace the heated breathing tube if the problem persists.
Wrong Tube	Med.	< 5 s	The heated breathing tube is not suitable for the selected therapy, or is damaged. Connect a suitable heated breathing tube. Replace the breathing tube if the problem persists.
Outlet Elbow Fault	Med.	< 5 s	A fault has been detected with the outlet elbow. Check that the outlet elbow is fully inserted into the Airvo 3 NIV. If the issue persists, replace the outlet elbow.
Outlet Elbow Too Warm	Med.	< 5 s	The outlet elbow is too warm to run the start up checks. Wait for the outlet elbow to cool down. If the issue persists, replace the outlet elbow.
Therapy alarm – high flow			
Chamber Leak Detected	Med.	< 30 s	The water chamber has been removed. Ensure the water chamber is correctly inserted into the Airvo 3 NIV. If the issue persists, contact your service representative.
Leak Detected	Med.	< 30 s	When used with Optiflow Junior 2 patient interfaces, the Airvo 3 NIV has detected a change in the leak of the breathing circuit. Check: - the water chamber has not been removed and is properly installed, - the heated breathing tube is plugged in correctly or is not damaged, - the patient interface is fitted correctly, and - the air filter is fitted correctly. If the issue persists, replace the consumables.
Blockage Detected	Med.	< 15 s	The Airvo 3 NIV has detected a blockage. Check: - for blockages in the heated breathing tube, patient interface and inlet air filter, - the patient interface is the correct size for the patient, and - the target flow rate is within the rated range of the interface. If the issue persists, replace the consumables.

Alarm condition	Priority	Delay	Meaning
Flow Below Target	Med.	< 2 min	The Airvo 3 NIV flow rate is lower than the target flow rate. Check: - for blockages in the heated breathing tube, patient interface and inlet air filter - the patient interface is the correct size for the patient, and - the target flow rate is within the rated range of the interface. If the issue persists, replace the consumables
Flow Above Target	Low	< 2 min	The Airvo 3 NIV flow rate is higher than the target flow rate. Check: - for leaks in the water chamber, heated breathing tube and patient interface, - the inlet air filter is inserted correctly, and - the target flow rate is within the rated range of the interface. If the issue persists, replace the consumables
Therapy alarm – NIV			
Blockage Detected	High	< 30 s	Air-path blocked resulting in degraded therapy and a CO₂ rebreathing risk. Raised when the leak (average flow) is less than 5 L/min and the RMS moving average of patient flow is less than 5 L/min for 15 seconds.
High Leak	High	< 30 s	Large leak in the system. Either the water chamber has been removed or the mask is off the patient. Raised when the filtered flow is greater than a threshold for more than 3 seconds.
Low Leak	High	< 30 s	There is little to no bias flow through the circuit which presents a CO₂ re-breathing risk. Raised when the leak (average flow) is less than 5 L/min and the root mean square (RMS) moving average of patient flow is greater than 5 L/min for 15 seconds.
High Inspiratory Pressure	High	3 breaths	Raised if the maximum estimated end of hose (EOH) pressure is above a user set limit during the inspiration portion of a breath.
Low Inspiratory Pressure	Med.	3 breaths	Raised if the maximum estimated EOH pressure is below a user set limit during the inspiration portion of a breath, for 3 consecutive breaths.
Therapy alarm – other			
Target Flow Too High	Med.	< 60 s	The Airvo 3 NIV has exceeded an internal temperature limit. Continued operation at the current settings may result in a device fault and reduced therapy. Check: - for blockages in the heated breathing tube, patient interface and inlet air filter, - the patient interface is the correct size for the patient, - the target flow rate is within the rated range of the interface, and - the ambient temperature is within the rated range of the device. This alarm will resolve when the internal temperature is within the expected range.
Check Water	Med.	< 30 min	The water chamber has run out of water. Replace the chamber and water bag to resume normal operation. Ensure that the water chamber and/or water bag are not allowed to run out of water to ensure continuous humidification of the breathing gases.
Humidity Below Target	Low	< 30 min	The Airvo 3 NIV cannot reach the target humidity. Check the water chamber contains water and the chamber base is not damaged. Consider reducing the target humidity or flow rate, if appropriate. If the issue persists, replace the water chamber.
Check Operating Conditions	Low	< 1 min	The Airvo 3 NIV has detected ambient conditions that are not suitable. Do not use the Airvo 3 NIV when the ambient temperature is below 18 °C or above 28 °C. Move the device to a suitable environment.

Alarm condition	Priority	Delay	Meaning
Oxygen alarms			
No O₂ Pressure at HPO Port	Med.	< 5 s	There is no oxygen being supplied to the high-pressure oxygen (HPO) inlet port during therapy. Check that the oxygen supply is working. If using an oxygen bottle, check the bottle is not empty. If switching to the low-pressure (LPO) inlet port or stopping oxygen delivery, set the FiO ₂ target to 21 %. Note: May switch to low-pressure oxygen (LPO) mode automatically if LPO is detected.
FiO₂ Below 25%	Med.	< 30 s	The oxygen being supplied to the LPO port has fallen below 25% during therapy. Check if the oxygen supply has been disconnected.
FiO₂ Below Target	Med.	< 2 min	The oxygen concentration being delivered is lower than the FiO ₂ target setting. Check the oxygen supply is properly connected to the HPO inlet port and there are no leaks at any oxygen hose connections. Make sure the number of connected devices does not exceed the capacity of the oxygen supply. Consider using the LPO connection if the oxygen supply has insufficient capacity.
FiO₂ Above Target	Med.	< 2 min	The oxygen concentration being delivered is higher than the FiO ₂ target setting. Check an oxygen supply is not connected to the low-pressure oxygen inlet port. Only one oxygen source should be used at a time. Check the oxygen supply is properly connected to the high-pressure oxygen inlet port and that there are no leaks at the oxygen hose connections.
High FiO₂ (LPO)	Med.	< 20 s	The FiO ₂ supplied by the LPO port is above the selected High Oxygen Alarm threshold for its intended clinical environment (range 30-95% or Off, default: 95%, see Airvo 3 NIV Technical Manual). Check FiO ₂ is appropriate for the patient's condition. Reduce FiO ₂ to the normal range when it is appropriate to do so.
Unexpected O₂	Med.	< 15 min	Oxygen is being supplied to the Airvo 3 NIV while in standby. Check all oxygen supplies are turned off and disconnected. If the issue persists, contact your service representative
High FiO₂ (HPO)	Med.	< 5 s	The FiO ₂ target is above the selected High Oxygen Alarm threshold for its intended clinical environment (range 30-95% or Off, default: 95%, see Airvo 3 NIV Technical Manual). Check FiO ₂ is appropriate for the patient's condition. Reduce FiO ₂ to the normal range when it is appropriate to do so.
Pulse oximetry alarms			
Pulse Ox Communication Failure	Med.	< 10 s	The Airvo 3 NIV is unable to communicate with the pulse oximeter. Check that the USB connector cable, sensor adapter cable and sensor cables are all properly connected. Replace the sensor cable, adapter cable then USB connector cable in turn if the problem persists.
Pulse Ox Not Recognized	Med.	< 10 s	The selected pulse oximeter has not been recognised. Please remove or change oximeter.
Pulse Ox Disconnected	Med.	< 1 s	The pulse oximetry accessories have become disconnected. Reconnect pulse oximetry accessories.
No Pulse Ox Sensor Connected	Med.	< 1 s	A pulse oximetry sensor cable was not detected or is inoperable. Check that the sensor cable is properly connected to the USB connector cable or replace the sensor cable if necessary.
Pulse Ox Sensor Off Patient	Med.	< 1 s	The pulse oximeter is no longer receiving SpO ₂ measurements from the patient. Check that the sensor is properly attached to a suitable measurement site on the patient.
No SpO₂ Reading	Med.	< 1 s (Masimo and Nellcor) < 16 s (Nonin)	The pulse oximeter is not sending valid SpO ₂ measurements. Check the sensor, cable and USB interface. Try replacing each component in turn until the problem is resolved.

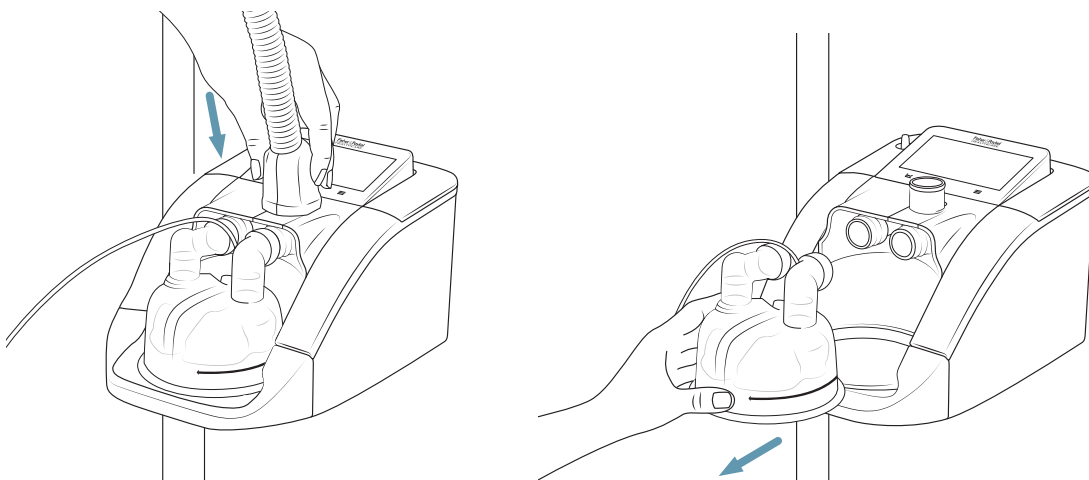
Alarm condition	Priority	Delay	Meaning
No Pulse Rate Reading	Med.	< 1 s (Masimo and Nellcor) < 16 s (Nonin)	The pulse oximeter is not sending valid Pulse Rate measurements. Check the sensor, cable and USB interface. Try replacing each component in turn until the problem is resolved.
Check Pulse Ox Cable/ Sensor	Med.	< 1 s	Masimo only. The pulse oximetry USB connector cable and/or the pulse oximetry sensor cable is not functioning correctly. Please remove and reconnect the accessory and if the problem persists replace the accessory.
Incompatible Pulse Ox Cable	Low	< 1 s	Masimo only. The pulse oximetry USB connector cable is incompatible. Please disconnect it from the device.
Incompatible Pulse Ox Sensor	Low	< 1 s	Masimo only. The pulse oximetry sensor cable is incompatible. Please disconnect it from the device.
Pulse Ox Cable Near Expiration	Low	< 1 s	Masimo only. The pulse oximetry USB connector cable is nearing expiration.
Pulse Ox Sensor Near Expiration	Low	< 1 s	Masimo only. The pulse oximetry sensor cable is nearing expiration.
Pulse Timeout	High	< 1 s	Nellcor only. The pulse oximetry USB connector cable is reporting the loss of a patients pulse signal. Check the condition of the patient.
SpO₂ Measurement Delayed	Low	< 1 s	Nellcor only. The pulse oximeter is indicating that its dynamic averaging time has exceeded 25 seconds for the SpO ₂ measurement.
Pulse Rate Measurement Delayed	Low	< 1 s	Nellcor only. The pulse oximeter is indicating that its dynamic averaging time has exceeded 25 seconds for the pulse rate measurement.
Pulse Ox Board Failure	Med.	< 1 s	Masimo and Nellcor only. The pulse oximetry USB connector cable has failed. Please remove and reconnect the cable and if the problem persists replace the cable.
Check Pulse Ox Sensor	Med.	< 1 s	Masimo and Nellcor only. The pulse oximetry sensor cable is not functioning correctly. Please remove and reconnect the cable and if the problem persists replace the cable.
Replace Pulse Ox Cable	Med.	< 1 s	Masimo and Nellcor only. There is a fault with the pulse oximetry USB connector cable and it needs replacement.
Replace Pulse Ox Sensor	Med.	< 1 s	Masimo and Nellcor only. There is a fault with the pulse oximetry sensor cable and it needs replacement.
Physiological alarms			
Low SpO₂	High	< 15 s	Check your patient. The SpO ₂ measurement has decreased below the Low SpO ₂ alarm threshold. Check that the alarm setting is appropriate for your patient (Range: 1-98%, default 85%, see Airvo 3 NIV Technical Manual).
High SpO₂	Med.	< 15 s	Check your patient. The SpO ₂ measurement has exceeded the High SpO ₂ alarm threshold. Check that the alarm setting is appropriate for your patient (Range: 2-99% or Off, default Off, see Airvo 3 NIV Technical Manual).
Low Respiratory Rate	Med.	3 breaths	Raised if the average respiratory rate is lower than the user set limit for 3 consecutive breaths. In Bi-Level S/T and Bi-Level PCV modes, if the Backup Rate is set equal to or higher than the Low Respiratory Rate the Low Respiratory Rate alarm is effectively disabled.
High Respiratory Rate	Med.	3 breaths	Raised if the average respiratory rate is higher than the user set limit for 3 consecutive breaths.
Low Minute Ventilation	Med.	3 breaths	Raised if the average expiratory minute ventilation is lower than the user set limit for 3 consecutive breaths.

Alarm condition	Priority	Delay	Meaning
Apnea Detected	Med.	User set	Apnea event detected. Raised if the time since the last inspiratory trigger (machine or spontaneous) is greater than the user set limit. In Bi-Level S/T and Bi-Level PCV modes, if the Backup Rate is set equal to or higher than (60/apnea) the apnea alarm is effectively disabled.
Disinfection alarms			
Disinfection Failed to Hold Temperature	Med.	< 3 min	The Airvo 3 NIV cannot heat up to the required disinfection temperature. Check: - the disinfection tube blue connector is connected to the top of the outlet elbow, - the disinfection tube red end is connected to the left hand chamber port, - the disinfection filter is connected to the right hand chamber port, Then restart the device. If the problem is not resolved, replace the disinfection tube and outlet elbow in turn. If the issue persists, contact your service representative.
Over Temperature Detected	Med	<5 s	The Airvo 3 NIV detected higher than expected temperatures during the disinfection cycle. Check: - the disinfection tube blue connector is connected to the top of the outlet elbow, - the disinfection tube red end is connected to the left hand chamber port, - the disinfection filter is connected to the right hand chamber port, Then restart the device. If the problem is not resolved, replace the disinfection tube and outlet elbow in turn. If the issue persists, contact your service representative.
Check Tube	Med.	<5 s	The Airvo 3 NIV cannot detect the disinfection tube. Check that the disinfection tube is not damaged and is plugged in correctly, then restart the device. If the problem is not resolved, replace the disinfection tube and outlet elbow in turn. If the issue persists, contact your service representative.
Leak Detected	Med.	<35 s	The Airvo 3 NIV has detected a leak in the disinfection circuit. Check: - the disinfection tube blue connector is connected to the top of the outlet elbow, - the disinfection tube red end is connected to the left hand chamber port, - the disinfection filter is connected to the right hand chamber port Then restart the device. If the problem is not resolved, replace the disinfection tube and outlet elbow in turn. If the issue persists, contact your service representative.
Blockage Detected	Med	<10 s	The Airvo 3 NIV has detected a blockage. Check that the disinfection tube is not blocked and that the disinfection filter is not wet, then restart the device. If the problem is not resolved, replace the disinfection tube and outlet elbow in turn. If the issue persists, contact your service representative.
Check Operating Conditions	Med.	< 1 min	The Airvo 3 NIV has detected ambient conditions that are not suitable. Do not use the Airvo 3 NIV when the ambient temperature is below 18 °C or above 28 °C. Move the device to a suitable environment, then restart the device. If the issue persists, contact your service representative
Wall Power Disconnected	Med.	< 5 s	The power cable has been removed and device is unable to perform a disinfection cycle. Connect device to wall power, then restart the device. If the issue persists, contact your service representative.
Unexpected O₂	Med.	< 1 min	Oxygen is being supplied to the Airvo 3 NIV while in disinfection mode. Check no oxygen is connected, then restart the device. If the issue persists, contact your service representative.

8. Reprocessing

Standard aseptic techniques should be followed to minimize contamination when handling the Airvo 3 NIV and accessories. The patient interface, heated breathing tube, water chamber and outlet elbow may become contaminated during use. As soon as possible after using the Airvo 3 NIV:

1. Remove the single-use accessories from the Airvo 3 NIV and dispose of them in accordance with local laws, regulations and hospital protocols for disposing of contaminated products.
 - Squeeze the sides of the breathing tube connector and lift to remove it from the Airvo 3 NIV.
 - Grip the port adapter and pull the water chamber away from the Airvo 3 NIV to remove it.



2. Reprocess the Airvo 3 NIV device exterior by following the instructions in section 8.1
3. Clean and high-level disinfect the Outlet Elbow by following the instructions in section 8.2.
4. Replace accessories within the maximum use period shown in section 8.3 (schedule for changing accessories).
5. Clean and disinfect pulse oximetry accessories (including reusable sensors) in accordance with the manufacturer's instructions.

Warnings

Do not clean and/or disinfect the Airvo 3 NIV while it is being used by a patient.

Do not submerge the Airvo 3 NIV or accessories in any cleaning solution or attempt to sterilize by autoclave, irradiation, steam, gas, ethylene oxide or any other method. This will seriously damage the device.

8.1 Airvo 3 NIV device exterior reprocessing

8.1.1 Clean

Equipment

- Mild detergent
- Clean, disposable, lint-free cloths
- Protective gloves

Cleaning Instructions

1. Mix a solution of warm water and mild detergent (refer to the detergent manufacturer's instructions for use).
2. Dampen a clean cloth with the cleaning solution.
3. Thoroughly wipe all outside surfaces of the device (including the Outlet Elbow) for at least one minute to remove any visible soil. Use the corner/edge of the cloth to clean all crevices of the device.

Rinse

4. Dampen a clean cloth with tap water.
5. Thoroughly wipe all outside surfaces of the device with the damp cloth to remove any disinfectant residue.

Dry

6. Thoroughly wipe all outside surfaces of the device with a dry cloth until it is visibly dry.
7. Allow to air dry until completely dry.

8.1.2 Disinfect**Perform disinfection only after all cleaning steps are complete****Equipment**

- Disinfectant wipes
- Clean, disposable, lint-free cloths
- Protective gloves

Disinfection Instructions

1. Use pre-soaked disinfecting wipes, to thoroughly wipe all outside surfaces of the device (including the Outlet Elbow).
2. Ensure that these surfaces remain visibly wet as directed by the manufacturer of the disinfecting wipes. Use additional wipes as needed to ensure that these surfaces remain wet for the required length of time.

Rinse

3. Dampen a clean cloth with tap water.
4. Thoroughly wipe all outside surfaces of the device with the damp cloth to remove any detergent residue.

Dry

5. Thoroughly wipe all outside surfaces of the device with a dry cloth until it is visibly dry.
6. Allow to air dry until completely dry.

⚠ Warnings

Other cleaning agents may be used if they are: non-abrasive, non-toxic, and non-corrosive. Do not use any cleaning agents that are not compatible with polycarbonate plastic.

Cleaning agents that are not suitable for use with the Airvo 3 NIV include: ammonia, ammonium hydroxide, caustic soda, iodine, methanol, methylated spirits, turpentine, and alkaline bleaches such as sodium hypochlorite. The use of any of these products will damage the Airvo 3 NIV.

Turn off and disconnect the Airvo 3 NIV from the wall power supply before cleaning to reduce the risk of electric shock.

Do not submerge the device in liquid of any kind.

Do not spray liquid directly onto the device.

Do not use rinse aids as these may cause damage to the outlet elbow.

These instructions have been validated by the manufacturer of the medical device as being capable of preparing a medical device for reuse. It remains the responsibility of the processor to ensure that the processing achieves the desired results, by using the correct equipment, materials, and personnel in the processing facility. This requires routine monitoring of the process.

8.2 Outlet elbow reprocessing

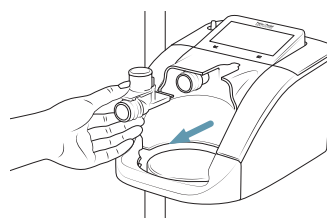
The Outlet Elbow requires cleaning and high-level disinfection. The Outlet Elbow can be reprocessed in two different ways:

- Disinfection Kit 900PT600 (see instructions in 900PT600)
- Washer-Disinfector (follow below instructions)

The outlet elbow can be removed from the Airvo 3 NIV for reprocessing by your central sterile services or reprocessing department. Reprocessing the outlet elbow must be performed in a washer-disinfector that complies with and is maintained, checked and validated to ANSI/AAMI ST15883-1:2009 (USA) and ISO 15883-1:2006 (outside USA).

Disassembly

Remove the outlet elbow from the Airvo 3 NIV. Firmly grab the rubber port-seal on the outlet elbow, push down on the grip lines with your thumb and pull the outlet elbow towards the front of the Airvo 3 NIV.



Transportation

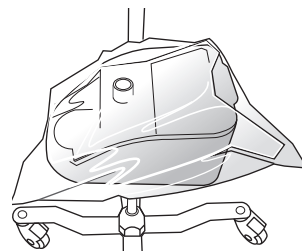
Follow hospital infection control protocols to package the outlet elbow for transport. Protect the outlet elbow from mechanical damage during transport.

Using the storage cover

It is important that the Airvo 3 NIV is stored properly after reprocessing. Store the Airvo 3 NIV in a clean, dry and dust-free location that is suitable for medical devices.

Cover the Airvo 3 NIV with the storage cover so that it remains clean during storage:

- Wrap the Airvo 3 NIV in a storage cover (900PT603) so that the identification label on the cover sits prominently above the display of the Airvo 3 NIV.
- Seal the cover with the adhesive tabs on the storage cover.



Cleaning and disinfection

Washer-disinfector supplies required for reprocessing of the Airvo 3 NIV outlet elbow are:

- Mildly alkaline cleaning agent such as neodisher® MediClean forte (0.2% v/v)

Place the outlet elbow in a washer-disinfector and orient the outlet elbow such that washing fluid can contact all internal surfaces and allow for draining. Run a cleaning and thermal high-level disinfection cycle:

- Pre-cleaning: Cold rinse for at least 1 minute
- Cleaning: Wash at 55 °C for at least 5 minutes with a mildly alkaline cleaning agent as per manufacturer's instructions (e.g. neodisher® MediClean forte, 0.2 % v/v)
- Neutralisation: Cold rinse for at least 1 minute
- Rinsing: Cold rinse for at least 1 minute
- Thermal disinfection: 90 °C for 5 minutes
- Drying: 90 °C for 25 minutes

! Note

Do not exceed the maximum use period for the outlet elbow.

Follow the manufacturer's instructions and warnings for all cleaning products.

Visual inspection

Visually inspect the outlet elbow for mechanical damage or discoloration of the chamber seal. If the seal or elbow appear damaged or discolored, replace the outlet elbow.

! Warning

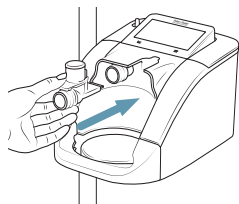
Do not use the outlet elbow if the seal or elbow appear damaged or discolored. A damaged outlet elbow may affect therapy delivery.

Storage and transport

It is important that the outlet elbow is stored properly after reprocessing. Store the outlet elbow in a clean, sealed plastic bag labeled with the disinfection process details. Follow your hospital protocol for storage of high-level disinfected devices. Protect the outlet elbow from mechanical damage during transport. Store the outlet elbow in a clean, dry and dust-free location that is suitable for medical devices. The outlet elbow can alternatively be inserted back into the Airvo 3 NIV then covered with the storage cover until the next use.

Reassembly

When setting up the Airvo 3 NIV for the next use follow the reassembly steps below. If reassembly occurs prior to the next use, cover the Airvo 3 NIV with the outlet elbow assembled with the clean storage cover.



Slide the disinfected outlet elbow into the slot above the chamber area on the Airvo 3 NIV.

Push firmly on the front of the elbow until the elbow locks into place.

! Note

Make sure the outlet elbow is installed in the Airvo 3 NIV before attaching the heated breathing tube.

8.3 Schedule for changing accessories

The Airvo 3 NIV accessories must be changed according to the schedule below. All single-patient-use accessories must be disposed of after the patient's therapy to prevent cross-contamination. Replace accessories within the period shown below, or immediately if they are damaged or discolored.

Accessory	Maximum Use
Optiflow Junior interfaces	1 week, or 1 patient (whichever comes first)
All NIV patient interfaces	Refer to instructions for use supplied with interface
Optiflow+ / Optiflow+ Duet interfaces Optiflow 3S interfaces All AirSpiral tube and chamber kits	14 days (7 days when using a nebuliser), or 1 patient (whichever comes first)
NIV tube and chamber kits (excluding NIV breathing tube filter, which must be changed more frequently)	14 days, or 1 patient (whichever comes first)
NIV breathing tube filter	24 hours or 1 patient (whichever comes first)
Air filter	3 months or 1000 hours use (whichever comes first)
Outlet elbow	5 years or 50 washer-disinfector cycles (whichever comes first)
Battery*	2 years from first use or 300 discharge cycles (whichever comes first)
Pulse oximetry accessories	Refer to instructions for use supplied with device.

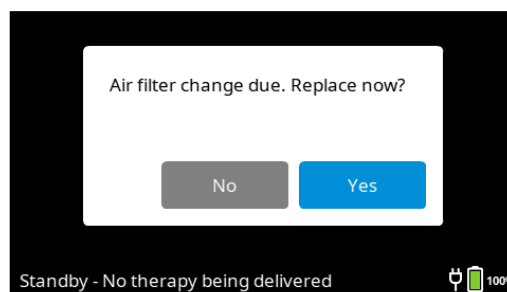
* See Airvo 3 Technical Manual for instructions to change the battery.

8.4 Replacing the air filter

The Airvo 3 NIV will display a message on startup when the air filter is due to be replaced.

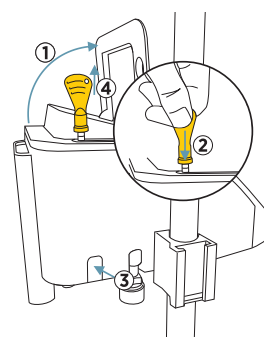
Begin by removing the old filter:

1. Raise the filter cover,
2. Push the filter removal tool down firmly onto the low-pressure oxygen inlet port to get the removal tool to grip,
3. Hold down the air-filter release button,
4. Pull up on the filter removal tool to remove the filter,
5. Insert the new filter and push down on top of the filter until it clicks into place,
6. Lower the filter cover.



8.5 Servicing

The Airvo 3 NIV does not require regular maintenance and contains no user serviceable parts. If the Medical Equipment system is modified from the specification of the manufacturer, evaluation to the requirements of 60601-1 standard is required. Refer to the Airvo 3 Technical Manual for product acceptance checks, functional tests and spare parts. Contact your Fisher & Paykel Healthcare representative if a fault develops or you are concerned the Airvo 3 NIV is not operating correctly



9. Pulse oximetry

The Airvo 3 NIV supports the following pulse oximeter accessories:

- The Masimo SET uSpO₂ Pulse Oximetry Cable (3412)
- The Medtronic Nellcor Oxicable (PMC10N-SF)
- The Nonin Xpod 3012LP USB (6703-001)
- The Nonin Xpod 3012HR USB Connector Cable (114403-001)

Pulse oximeters primarily measure the oxygen saturation of blood and pulse rate. Oxygen saturation or SpO₂ is the percentage of hemoglobin in the blood that is saturated with oxygen. Hemoglobin is a protein in the red blood cells that carries oxygen from the lungs to the rest of the body. Pulse rate is the number of heart beats per minute.

A pulse oximeter operates by emitting two different wavelengths of light, typically red and infrared, into a perfused tissue site such as a fingertip or earlobe. The emitted light passes through the tissue, and a photodetector in the sensor measures the intensity of the transmitted or reflected light. By comparing the absorption of red and infrared light, the pulse oximeter calculates the ratio of oxygenated to total hemoglobin in the blood, which is used to determine the oxygen saturation level. Additionally, the device analyzes the variations in the light intensity to measure the patient's pulse rate.

9.1 Pulse oximetry warnings, cautions, and notes

Warnings

- In line with the indications for use of the Airvo 3 NIV, the monitoring functionality of the Airvo 3 NIV, is intended for use on spontaneously breathing patients and not intended for patients requiring life support. It is the responsibility of the clinician to select the appropriate level of monitoring for their patient and to be prepared to deal with alarms and equipment malfunction. Additional, independent monitoring equipment may be necessary.
- If any measurement seems questionable, first check the patient's vital signs by alternate means and then check the pulse oximeter for proper functioning.
- Using different alarm settings on devices within a single area, such as an intensive care unit, can cause a hazard.
- Periodically reposition the sensor to help prevent ischemia.
- If any measurement seems questionable, first check the patient's vital signs by alternate means. Then check the pulse oximeter USB connector, adapter, sensor, and Airvo 3 NIV for proper operation.
- Carefully route cabling to reduce the possibility of patient entanglement or strangulation.
- Do not use single-patient-use pulse oximeter sensors on more than one patient to avoid cross-infection and/or contamination.
- Refer to the third party pulse oximeter sensor instructions for information on the risks of re-using a single use sensor.
- Follow the user instructions supplied with multi-use pulse oximeter sensors, adapters and USB connector cables to clean and disinfect these devices between patients to avoid cross-infection and/or contamination.
- Tissue damage may be caused by incorrect application of the sensor, e.g. by wrapping the sensor too tightly. Follow the instructions supplied with the sensor for correct application.
- Use only compatible oximetry sensors and accessories for SpO₂ and pulse rate measurements. Verify compatibility before use to avoid incorrect operation of your Airvo 3 NIV, inaccurate measurements and/or patient injury. See Appendix 3 for a list of compatible accessories.
- Supplemental oxygen will attenuate patterns of desaturation. A patient's respiratory compromise can be proportionally more severe before patterns appear in the saturation trend. Remain vigilant when monitoring a patient on supplemental oxygen.
- Explosive hazard: Do not use this device in an explosive atmosphere or in the presence of flammable anesthetics or gases.
- To protect from electric shock, always remove the sensor and completely disconnect the pulse oximeter before bathing the patient.

Nellcor:

- Do not rely solely on the Nellcor Oxicable for patient assessment. It must be used in conjunction with clinical signs and symptoms.
- Avoid spilling liquids into the Nellcor Oxicable.
- Refer to the Oxicable PMC10N-SF instructions for use for a complete list of warnings and more information when using the Medtronic Oxicable with the Airvo 3 NIV.

Nonin:

- Operation of the Nonin Xpod USB connector below the minimum amplitude of 0.3% modulation may cause inaccurate results.

Cautions

- Before cleaning the pulse oximetry accessories, disconnect the device from the Airvo 3 NIV to avoid electrical shock and flammability hazards.
- Do not place the pulse oximetry accessories on electrical equipment that may affect the device, preventing it from working properly.
- To minimize radio interference, other electrical equipment that emits radio frequency transmissions should not be in close proximity to the pulse oximeter equipment.
- Do not place the Airvo 3 NIV where controls can be changed by the patient.
- The accuracy of the SpO₂ measurement may be affected if the total sensor cable length (including extension cables) is greater than 3 meters.
- Setting extreme alarm thresholds can render alarms useless and may lead to patient injury.

- Inspect the sensor application site at least every 6 to 8 hours to ensure correct sensor alignment and skin integrity. Patient sensitivity may vary due to medical status or skin condition. Discontinue use of adhesive tape sensors if the patient exhibits an allergic reaction to the adhesive material.

- Check that the pulse oximetry alarm limits are appropriate for the patient being monitored every time pulse oximetry is used.

Masimo:

- If the Low Perfusion message is frequently displayed, find a better perfused monitoring site. In the interim, assess the patient and, if indicated, verify oxygenation status through other means.
- Change the application site or replace the sensor and/or patient cable when a “Replace Pulse Ox Sensor” and/or “Replace Pulse Ox Cable”, or a persistent poor signal quality message (such as “Low Signal I.Q.”) is displayed on the Airvo 3 NIV. These messages may indicate that patient monitoring time is exhausted on the patient cable or sensor.
- If using pulse oximetry during full body irradiation, keep the sensor out of the radiation field. If the sensor is exposed to the radiation, the reading might be inaccurate or the device might read zero for the duration of the active irradiation period.
- Variation in measurements may be profound and may be affected by sampling technique as well as the patient's physiological conditions. Any results exhibiting inconsistency with the patient's clinical status should be repeated and/or supplemented with additional test data. Blood samples should be analyzed by laboratory instruments prior to clinical decision making to completely understand the patient's condition.
- Replace the cable or sensor when a “replace sensor” or when a “low SIQ” message is consistently displayed while monitoring consecutive patients after completing the troubleshooting steps listed in this manual.
- Please refer to the instructions for use provided with the Masimo uSpO₂ Pulse Oximetry Cable for additional warnings, cautions, and notes.
- The Masimo SET uSpO₂ Pulse Oximetry Cable (Oximetry Cable) must be connected to the Airvo 3 NIV for power, communication, display and alarm management.

Nonin:

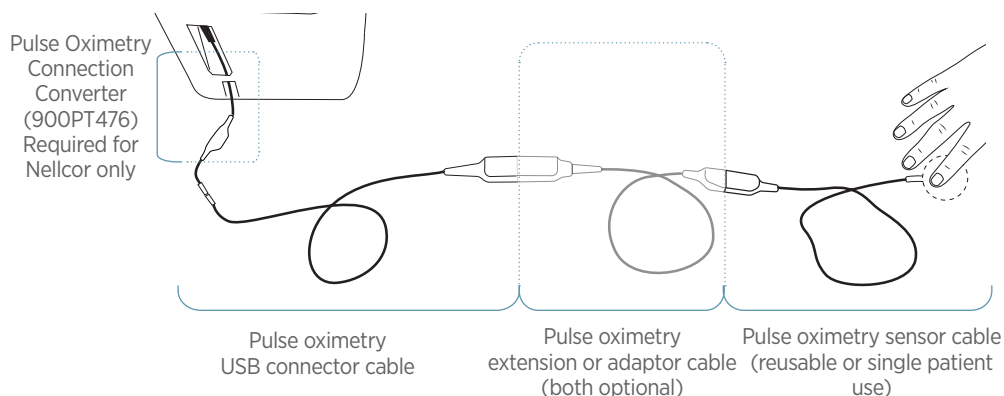
- The Nonin Xpod USB connector has motion tolerant software that minimizes the likelihood of motion artefact being misinterpreted as good pulse quality. In some circumstances, however, this device may still interpret motion as good pulse quality. This covers all available outputs (i.e. SpO₂, PR, PLETH, PPG).

! Note

- The accuracy of pulse oximeter sensors, adapters and patient cables cannot be assessed using a functional tester.
- High-intensity extreme lights (such as pulsating strobe lights) directed on the sensor, may not allow the pulse oximeter to obtain vital sign readings.
- When using the Maximum Sensitivity setting, performance of the “Sensor Off” detection may be compromised. If the device is in this setting and the sensor becomes dislodged from the patient, the potential for false readings may occur due to environmental “noise” such as light, vibration, and excessive air movement.
- For more information about required safety and regulatory requirements for pulse oximeters, refer to ISO 80601-2-61, and IEC 60601-1. Additional safety information can be found in the labelling provided with each sensor.
- Do not loop the pulse oximetry cabling into a tight coil or wrap around the device, as this can damage the patient cabling.
- Additional information specific to the sensors compatible with the Airvo 3 NIV pulse oximeter, including information about parameter/measurement performance during motion and low perfusion, may be found in the sensor's directions for use.
- Cables and sensors are provided with X-Cal™ technology to minimize the risk of inaccurate readings and unanticipated loss of patient monitoring. Refer to the Cable or Sensor directions for use for the specified duration of the patient monitoring time.
- Each specific sensor comes with manufacturer-supplied instructions for its use. Please refer to these for further details, including Bland Altman plots.

9.2 Setup for pulse oximetry

Connect the pulse oximetry USB connector cable to either USB port on the back of the Airvo 3 NIV. Clip the cable into the cable tidy so that it is not pulled out accidentally. The Airvo 3 NIV will automatically detect a compatible pulse oximeter.



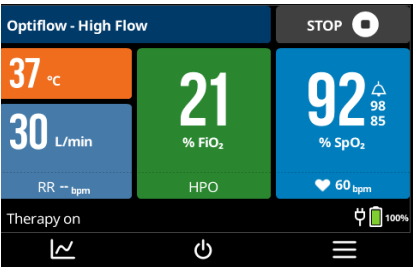
9.2.1 Pulse oximetry accessories

When compatible pulse oximetry accessories are connected, the Airvo 3 NIV can display:

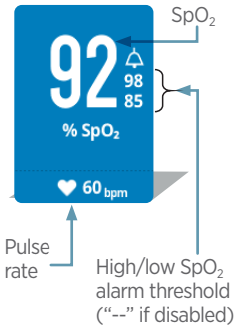
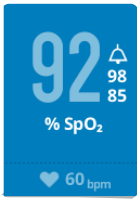
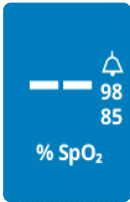
- functional oxygen saturation (no pulse rate alarms are included in the Airvo 3 NIV),
- pulse rate (no pulse rate alarms are provided),
- perfusion index (Masimo only),
- plethysmograph, and
- signal quality indicators.

9.3 During therapy

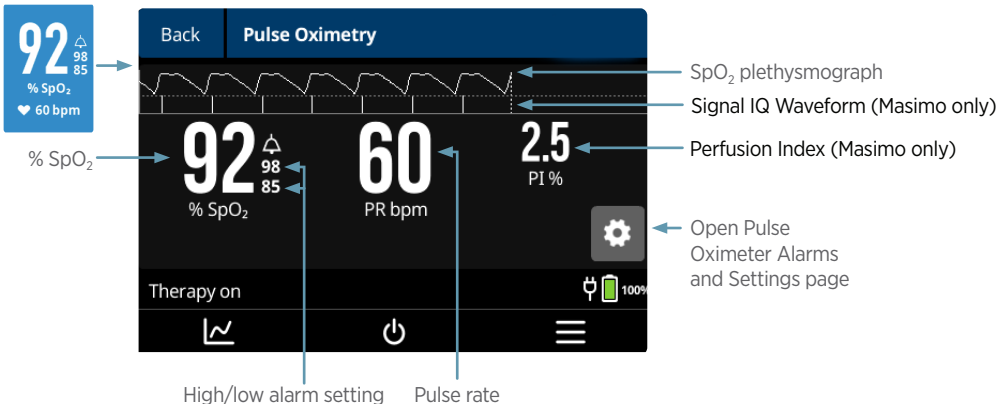
The Pulse Oximetry tile will be automatically displayed on the Home screen when a compatible pulse oximetry USB connector cable is connected to the Airvo 3 NIV.

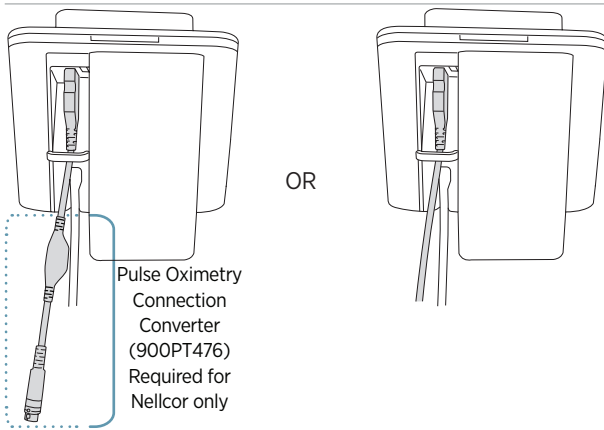


Pulse oximetry measurements and status are shown as follows:



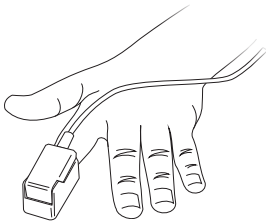
Tap the Pulse Oximetry tile to open the Pulse Oximetry screen.





Connect sensor to Airvo 3 NIV

Connect the pulse oximeter USB connector cable to the USB port on the back of the Airvo 3 NIV. The Airvo 3 NIV will automatically detect compatible devices. Connect the pulse oximetry sensor cable into the USB connector cable.



Attach sensor to patient

Carefully select a pulse oximetry sensor based on the patient's age, weight and intended sensor application site. More information can be found in the instructions supplied with each sensor.

⚠ Warnings

Inaccurate SpO₂ and/or pulse-rate readings may be caused by:

- Improper sensor application and placement.
- Elevated levels of Carboxyhemoglobin (COHb) or methaemoglobin (MetHb): High levels of COHb or MetHb may occur with a seemingly normal SpO₂. When elevated levels of COHb or MetHb are suspected, laboratory analysis (COOximetry) of a blood sample should be performed.
- Elevated levels of bilirubin.
- Elevated levels of dyshemoglobin.
- Vasospastic disease, such as Raynaud's, and peripheral vascular disease.
- Hemoglobinopathies and synthesis disorders such as thalassemia, Hb s, Hb c, and sickle cell, etc.
- Hypocapnic or hypercapnic conditions.
- Severe anemia.
- Very low arterial perfusion.
- Extreme motion artifact.
- Abnormal venous pulsation or constriction.
- Severe vasoconstriction or hypothermia.
- Arterial catheters and intra-aortic balloon.
- Intravascular dyes, such as indocyanine green or methylene blue.
- Externally applied coloring and texture, such as nail polish, acrylic nails, glitter, etc.
- Birthmark(s), tattoos, skin discolorations, moisture on skin, deformed or abnormal fingers etc.
- Skin color disorders.
- Excessive ambient light.
- Excessive motion.
- Electrosurgical interference.
- Blood flow restrictors (arterial catheters, blood pressure cuffs, infusion lines, etc.).
- Moisture in the sensor.
- Improperly applied sensor.
- Incorrect sensor type.
- Poor pulse quality.
- Venous pulsations.
- Low haemoglobin concentrations.
- A sensor not at heart level.

Dyes or any substance containing dyes that change usual blood pigmentation may cause erroneous readings.

Loss of monitoring can result if any objects hinder the pulse measurement. Ensure that no blood flow restrictors (e.g., blood pressure cuff) hinder pulse measurements.

The oximeter sensor may not work on cold extremities due to reduced circulation. Warm or rub the finger to increase circulation or reposition the sensor.

Inaccurate readings can result due to residue (e.g. dried blood) in light path or degradation of optical characteristics of sensor components. Refer to cleaning instructions supplied with the pulse oximetry accessories.

False high readings can result if SpO₂ is low due to dysfunctional hemoglobin (e.g. carboxyhemoglobin or methemoglobin).

Read the instructions supplied with the pulse oximetry accessories for additional safety information (including any potential risks or adverse effects from sensor materials), measurement site selection, detailed sensor setup, maximum sensor application time at a single site before repositioning, cable lifetime, sensor lifetime, factors that can interfere with measurement, troubleshooting and maintenance instructions.

9.4 Description of measurements

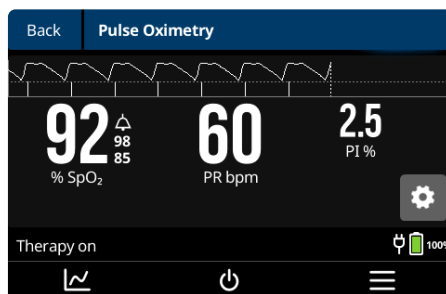
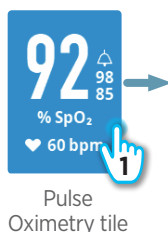
Pulse oximetry measurements are displayed on the Pulse Oximetry tile, the Pulse Oximetry screen and the Data and Graphs screen. Measurements are updated every second.

Tap the Pulse Oximetry tile to open the Pulse Oximetry screen and  to open the Data and Graphs screen.

Tapping  on the Pulse Oximetry screen provides a shortcut to the pulse oximetry alarms and settings.

Caution

If any measurement seems questionable, check the patient's vital signs by another method. Then check the pulse oximetry accessories and Airvo 3 NIV are set up, configured and working correctly.



Pulse Oximetry screen

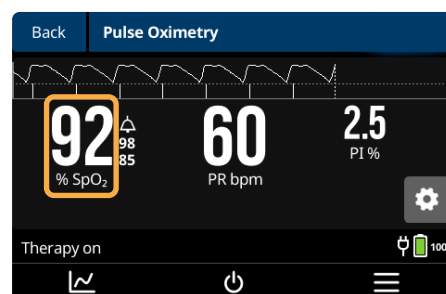
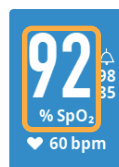
9.4.1 SpO₂

The Airvo 3 NIV is calibrated to display functional oxygen saturation (SpO₂) as a percentage. The SpO₂ value displayed is an average of measurements over a user selectable period (see Averaging Time in section 9.5 below). A long averaging period will generally produce more stable values but the SpO₂ displayed will respond more slowly to rapid changes in arterial blood oxygen saturation (SaO₂).

Stability of the SpO₂ measurements displayed may provide a good indication of a valid signal. Though stability is a relative term, experience with the device and patient observations will help you separate physiological effects from artifacts caused by a poorly-placed sensor or excess patient motion.

Inconsistencies between SpO₂ displayed on the Airvo 3 NIV and arterial blood gas analysis or clinical assessment may be caused by:

- poor signal quality,
- low perfusion,
- improperly-placed sensors or cables, and/or
- the patient's condition.

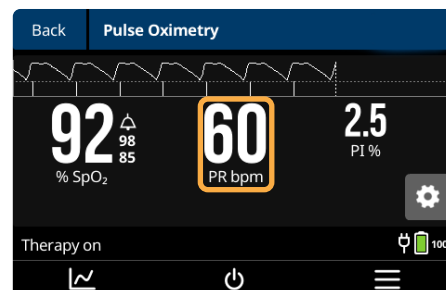
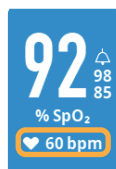


9.4.2 Pulse rate

Pulse rate (, PR) measurements are based on optical detection of pulsatile peripheral blood flow by the pulse oximetry sensor. The pulse rate displayed, in beats per minute (bpm), is an average of measurements over a user-selectable period.

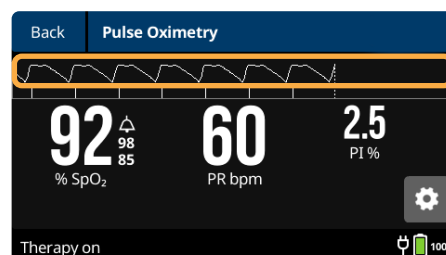
Small differences in the pulse rate displayed by different equipment may be caused by different approaches to averaging. Small discrepancies between cardiac electrical activity and pulse rate obtained from peripheral measurements can also arise. Large discrepancies between equipment may be caused by:

- poor signal quality,
- low perfusion,
- improperly placed sensors or cables, and/or
- the patient's condition.



9.4.3 Plethysmograph

A plethysmograph (or photo-plethysmograph) provides a non-normalized indication of the change in blood volume measured by the pulse oximeter sensor. The shape of the plethysmograph may change between patients, between measurement sites and for different sensor models. The plethysmograph provides an indication of signal inadequacy. A low amplitude or variable plethysmograph may indicate poor/inadequate signal. The plethysmograph is displayed on the Pulse Oximetry screen.



9.4.4 Signal IQ Waveform (Masimo)

The Signal IQ waveform shows the measurement confidence and timing of each detected pulse relative to the plethysmograph waveform. The height of the vertical bars indicate the relative confidence of the measurement. A high bar corresponds to higher confidence. In addition, the bars should visually correlate to the peak of the plethysmograph waveform. This along with perfusion index provides a better tool for assessing potential problems such as blood flow obstructions, poor sensor placement, artifacts or interference.

If Signal IQ is low an indication will occur with the status message "Low SpO₂ Signal IQ" displayed. During this time the SpO₂ and pulse rate numbers will be greyed out.

9.4.5 Perfusion Index (Masimo)

The Perfusion Index (PI) is the ratio of the pulsatile blood flow to the non-pulsatile or static blood in peripheral tissue. Perfusion index represents a noninvasive measure of peripheral perfusion that can be continuously and noninvasively obtained from a pulse oximeter.

9.4.6 Signal quality indicators (Nonin)

Nonin pulse oximetry equipment indicate signal quality based on the perfusion of the patient. There are three states: green, yellow, and red corresponding to high, low/marginal, and low/poor signal quality respectively. The Airvo 3 NIV indicates low signal quality by greying out the SpO₂ and Pulse rate numbers.

9.5 Description of settings and alarms

This section describes the behavior of pulse oximetry settings and alarms. See the Alarms and measurement section (9.6) on how to make changes to the alarm thresholds and settings.

9.5.1 Patient alarm thresholds

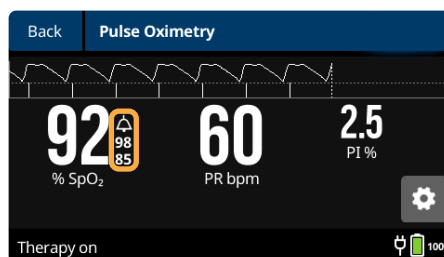
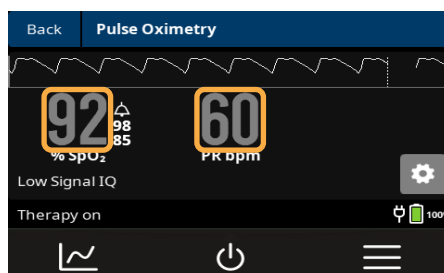
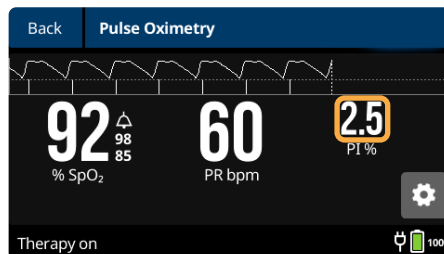
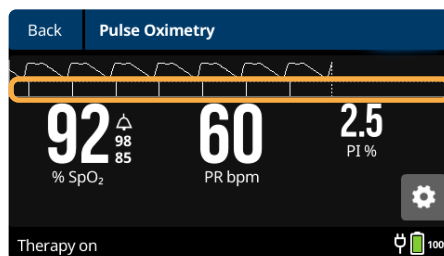
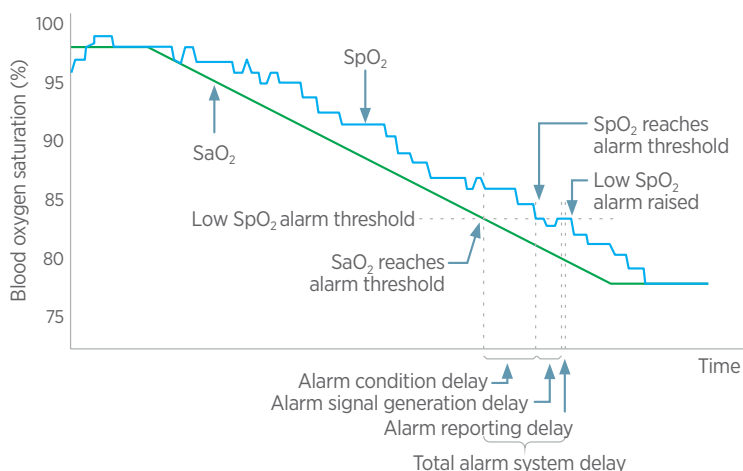
The following alarms can alert you to changes in your patient's condition:

- SpO₂ Low alarm
- SpO₂ High alarm

The corresponding alarm will be raised when a measurement is lower or higher than the alarm threshold. SpO₂ alarm thresholds are displayed on the Pulse Oximeter tile and Pulse oximeter screen.

9.5.2 SpO₂ Alarm Delay

The SpO₂ Alarm Delay setting defers the Low SpO₂ and High SpO₂ alarms for up to 15 seconds. This delay helps reduce non-actionable alarms for short desaturations. The alarm will start, if the alarm condition remains, after the delay.



9.5.3 Alarm response time

Audible and visual alarms are subject to an alarm response delay. The alarm response delay has three parts, the:

1. Alarm condition delay: the duration for a physiological change to be recognized by a pulse oximeter,
 2. Alarm signal generation delay: the period between detecting a condition and signaling the alarm, and
 3. Alarm reporting delay: the period between receiving an alarm signal from a monitoring device and reporting the alarm to the user.
- Measurement averaging will affect the signal generation delay: a larger averaging time will increase the signal generation delay. These delay concepts are illustrated on the graph below for a decrease in SaO_2 leading to a SpO_2 Low alarm as an example. The illustration does not reflect the actual length of delays. Refer to ISO 80601-2-61 for more information about alarm response delay.

9.5.4 Averaging time

The SpO_2 averaging time can be adjusted depending on patient acuity and area of care. This is the time, in seconds for Masimo, and heart beats for Nonin that measurements are averaged over. Shorter averaging times are sometimes preferred in, for example, sleep testing while longer averaging times are more suited to telemetry and neonates.

9.5.5 Sensitivity modes (Masimo)

There are three sensitivity modes. Normal sensitivity is recommended for patients who are experiencing some compromise in blood flow or perfusion. These patients are usually observed frequently such as in the intensive care unit. Adaptive Probe Off Detection (APOD) sensitivity is recommended when there is a higher probability of the sensor becoming detached. It is also the recommended mode for areas where patients are not visually monitored continuously. It provides enhanced protection against erroneous pulse rate and SpO_2 readings when the sensor becomes inadvertently detached from a patient due to excessive movement. Maximum sensitivity (MAX) is recommended for patients with weak signals, useful during procedures or when clinician contact is continuous such as high acuity areas.

9.5.6 Response mode (Nellcor)

The Nellcor Oxicable provides a Normal and Fast mode response modes. In Normal Mode, the SpO_2 averaging time is six to seven seconds and in Fast Mode, the SpO_2 averaging time is approximately three seconds. Pulse rate averaging time is approximately five seconds, independent of response mode.

9.6 Alarm and measurement settings

To change pulse oximetry alarm thresholds and settings:

1. Tap  to open the system menu,
2. Select Pulse Oximeter Alarms and Settings,
3. Tap the desired setting, scrolling if necessary,
4. Use the + / - buttons to select the required value,
5. Tap Confirm to apply the change or Cancel to discard any changes and return to the settings list.

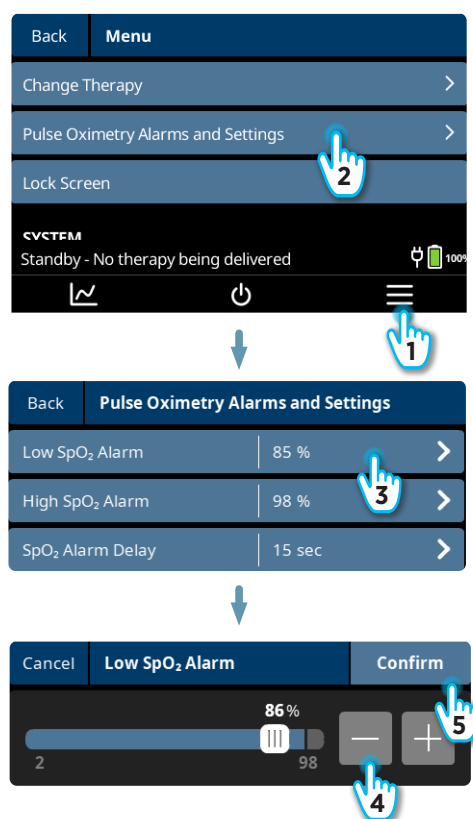
Tap Back twice to return to the Home Screen when you have finished making changes.

All settings are persistent and will retain their previous value when the Airvo 3 NIV is turned on and Same Patient is selected. Selecting New Patient when reviewing the disinfection state applies the default values for its intended clinical environment to all alarm and measurement settings.

Refer to the troubleshooting section for troubleshooting SpO_2 measurements and general device alarms.

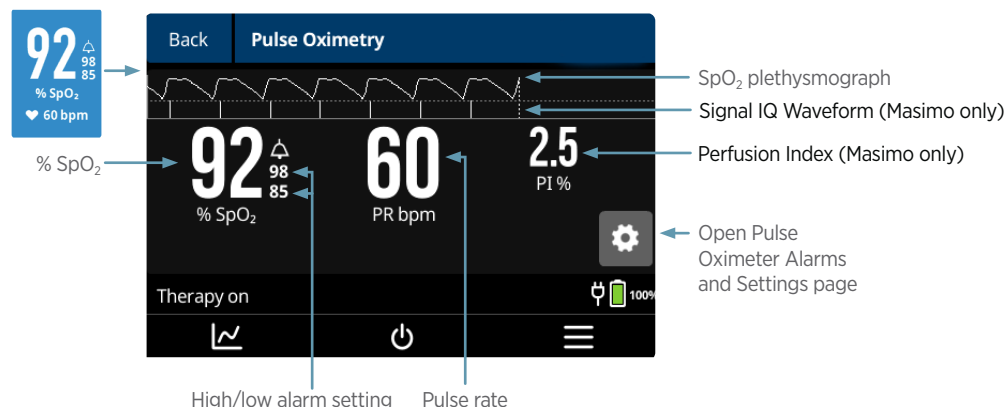
Label	Description	Factory default	Range
Low SpO_2 Alarm^{1,2}	Threshold for SpO_2 Low alarm	85%	1 – 98% ³
High SpO_2 Alarm²	Threshold for SpO_2 High alarm	Off	Off, 2 – 99% ³
SpO_2 Alarm Delay	Delay before audible Low SpO_2 or High SpO_2 alarm	15 seconds	0, 5, 10, 15 seconds
Averaging Time	Masimo: The length of time to average over Nonin: The number of pulses to average over	8 seconds 8 beats	2-4, 4-6, 8, 10, 12, 14, 16 seconds 4 or 8 beats
Sensitivity Mode	Masimo Only	APOD	Normal, APOD, Max
Response Mode	Nellcor Only	Normal	Normal, Fast

¹ The minimum threshold may be set when the device is set up for its intended clinical environment. Refer to the Airvo 3 Technical Manual for details.



2. The high alarm threshold cannot be set below the low alarm threshold.
3. The alarm threshold can be changed in 1% steps.
4. Max sensitivity mode does not persist through an Airvo 3 power cycle power cycle. Once powered back up the sensitivity setting reverts to the current default setting.

9.7 Troubleshooting



To help ensure successful monitoring of your patient's SpO₂:

- Apply the pulse oximeter sensor to a well-perfused site.
- Select a measurement site that has unrestricted blood flow.
- Follow all the instructions supplied with the pulse oximetry sensor to ensure the device is correctly applied.

The Pulse oximeter status bar displays the status of the pulse oximeter. Tap the Pulse oximeter tile to open the Pulse Oximetry screen and view the status. Possible status messages and warnings are described below.

Message	Cause/remedy
Low SpO₂ Signal IQ	Masimo only. The pulse oximeter is indicating low signal confidence in the values displayed due to poor signal strength. The parameters displayed are greyed out while in this state. The patient should be assessed and the sensor should be checked for correct application.
Low Signal quality	Nonin only. The pulse oximeter is indicating low signal quality. The parameters are displayed in gray while in this state. These low signal quality states may be caused by excess motion, low perfusion, a long/blocked light path, or a damaged or incorrectly fitted sensor. • Follow the sensor's user instructions to check it is the correct type and that it has been correctly applied to the patient. • Reduce or eliminate motion at the monitoring site. • Consider an adhesive sensor. • Check that the sensor's emitter and detector are properly aligned, particularly when using an adhesive sensor. • Consider a different measurement site. • Check that blood flow to the measurement site is not restricted. • See the pulse oximetry section for physiological conditions that may affect pulse oximetry measurement accuracy and consider an alternative method if indicated. • Remove excessive fingernail polish or artificial nails. • Replace the sensor.
Replace Cable Next Patient	Masimo only. The pulse oximetry USB connector cable is defective or has expired and should be replaced after the current patient.
Replace Sensor Next Patient	Masimo only. The pulse oximetry sensor cable is defective or has expired and should be replaced after the current patient.
Sensor Initialising	Masimo only. The pulse oximetry sensor is initialising. If values are not displayed within 30 seconds disconnect and reconnect the sensor. If the problem persists replace the sensor.

Message	Cause/remedy
Low Perfusion Index	Masimo only. The pulse oximeter is indicating that the perfusion index of the patient is low. Please move the sensor to a better perfused site.
Demo mode	Masimo only. The pulse oximeter is indicating that it is running in demonstration mode. If this is unintentional please remove and replace the oximetry equipment.
SpO₂ Only Mode	Masimo only. The pulse oximeter is indicating that it is running in SpO ₂ only mode. No pulse rate is available. Check the sensor placement referring to the directions for use provided with the sensor.
Pulse Search	Masimo and Nellcor only. The pulse oximeter is performing a pulse search. For Nellcor a continuous message indicates a loss-of-pulse while a flashing message indicates searching. If values are not displayed within 30 seconds, disconnect and reconnect the sensor. If the problem persists replace the sensor.
Interference Detected	Masimo and Nellcor only. The pulse oximeter is indicating that interference has been detected. Check that the sensor is correctly applied and if necessary cover the sensor site with an opaque material.
Patient Missing	The pulse oximeter cannot detect a patient. Check that the sensor is properly fitted by following the user instructions supplied with the sensor.
Sensor Disconnected	The pulse oximeter is indicating that there is no sensor currently connected. If there is remove and reconnect. If the problem persists please replace the oximeter accessories.

Nellcor recommends the following actions if there are any issues with their oximetry:

- Reposition sensor,
- Ensure the sensor is not too tight,
- Try an alternative sensor placement sight,
- Optically cover the sensor sight,
- Use an adhesive sensor,
- Use an ear, nasal, or forehead sensor,
- Use a headband with the forehead sensor,
- Check the assembly,
- Remove any nail polish from nail beds,
- Check for and eliminate external interference,
- Clean the sensor site,
- Secure the sensor cable.

If pulse oximeter measurements do not correlate with clinical assessment and/or arterial blood gas measurements:

- check the pulse oximeter status, as described above,
- check the pulse oximeter sensor is fitted correctly, following the user instructions supplied with the sensor,
- review the pulse oximetry section for conditions that may affect pulse oximetry measurement accuracy and consider an alternative method if indicated, and/or
- try a different measurement site.

If the Airvo 3 NIV loses mains power, the system will automatically switch over to use the internal battery and pulse oximetry functionality will be maintained including patient settings and trend data.

If the Airvo 3 NIV loses mains power and the battery is depleted, pulse oximetry functionality will be lost. It will be restored once power is restored to Airvo 3 NIV maintaining patient settings but trend data will be lost.

10. Adult NIV therapy

This section describes noninvasive ventilation (NIV) therapy delivery and settings. Refer to sections 4 and 5 to set up and use the Airvo 3 NIV to deliver NIV therapy.

10.1 Therapy description

The Airvo 3 NIV provides NIV therapy using a single-limb circuit and a sealed nasal, oronasal or full-face patient interface. Three NIV modes are supported:

- 1. CPAP (continuous positive airway pressure),
- 2. Bi-level S/T (spontaneous/timed), and
- 3. Bi-level PCV (pressure-controlled ventilation).

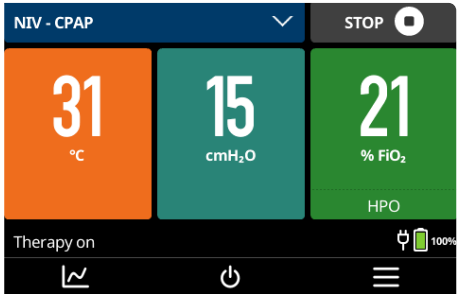
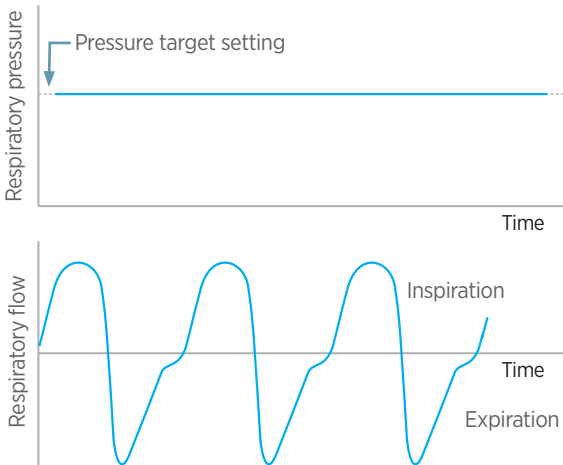
The main differences between these modes are the characteristics of spontaneous and machine breaths (see table below). Supplementary oxygen (if required) is blended with room air to increase the concentration of oxygen in the respiratory gases supplied.

The source for triggering and timing of spontaneous and machine breaths is shown for each of the NIV modes supported.

Spontaneous breaths			Machine breaths	
Mode	Trigger	Inspiration duration	Trigger	Inspiration duration
CPAP	n/a	Patient	n/a	n/a
Bi-level S/T	Patient	Patient	Backup rate	Inspiration time
Bi-level PCV	Patient	Inspiration time	Backup rate	Inspiration time

10.1.1 CPAP mode

In CPAP mode, the Airvo 3 NIV supplies breathing gases to the patient at a constant airway pressure. Airway pressure in the sealed mask is created by adjusting the flow generator automatically to maintain the target CPAP setting. The target CPAP is set to the prescribed value by the operator.



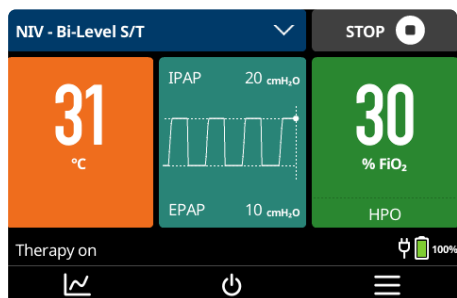
The Airvo 3 NIV includes a pressure ramp function for CPAP mode, which may help your patient acclimatize to NIV therapy by temporarily reducing CPAP. See section 11.1.8 for information on using the ramp function.

10.1.2 Bi-level S/T mode

In Bi-level S/T mode, the Airvo 3 NIV supports the patient's respiration by increasing the airway pressure during inhalation and decreasing the airway pressure during exhalation. Airway pressure is inspiratory positive airway pressure (IPAP) during inhalation and expiratory positive airway pressure (EPAP) during exhalation. Both values are set to prescribed values by the operator.

Generally, the timing and duration of breaths are controlled by the patient: airway pressure is increased when a breath is triggered and decreased when the breath cycles. Patient triggered breaths are limited to a maximum duration of 3 seconds.

The Airvo 3 NIV includes a pressure ramp function for Bi-level S/T mode, which may help your patient acclimatize to NIV therapy by temporarily reducing IPAP/EPAP. See section 11.1.8 for more information on the ramp function.

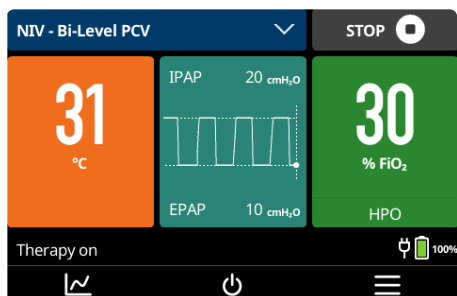


10.1.3 Bi-level PCV mode

In Bi-level PCV mode, the Airvo 3 NIV supports the patient's respiration by providing IPAP for a fixed amount of time after detecting each inhalation, then returning to EPAP. Both values are set to prescribed values by the operator.

Generally, the start of breaths is controlled by the patient: airway pressure is increased when a breath is triggered, but the Airvo 3 NIV controls the duration of inspiration using the Inspiration Time setting.

The Airvo 3 NIV includes a pressure ramp function for Bi-level PCV mode, which may help your patient acclimatize to NIV therapy by temporarily reducing IPAP/EPAP.



10.1.4 Backup breaths

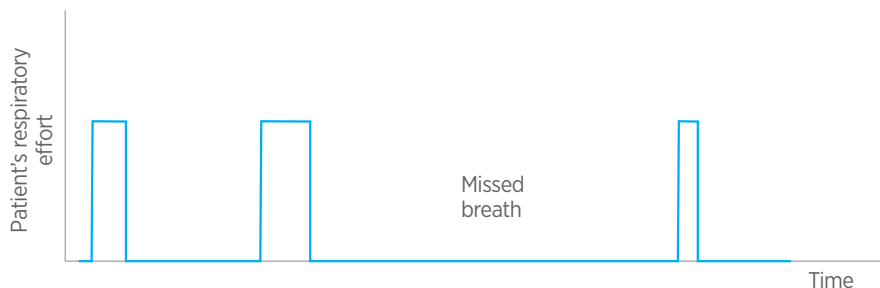
The Airvo 3 NIV will automatically supply backup breaths when breaths are not triggered by the patient in Bi-level S/T and Bi-level PCV modes. Backup breaths do not apply to CPAP mode.

Backup breaths are supplied whenever the patient's breathing rate falls below the Backup Rate setting. The length of inspiration for each backup breath is set by the Inspiration Time.

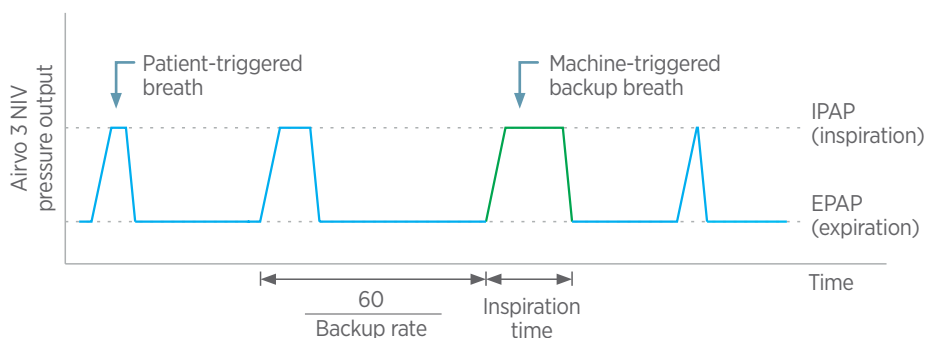
10.1.5 Comparing Bi-level S/T and Bi-level PCV modes

In both Bi-level S/T and Bi-level PCV modes, the airway pressure increases from EPAP to IPAP when the patient triggers a breath. The main difference between these two modes is the duration of the inhalation phase.

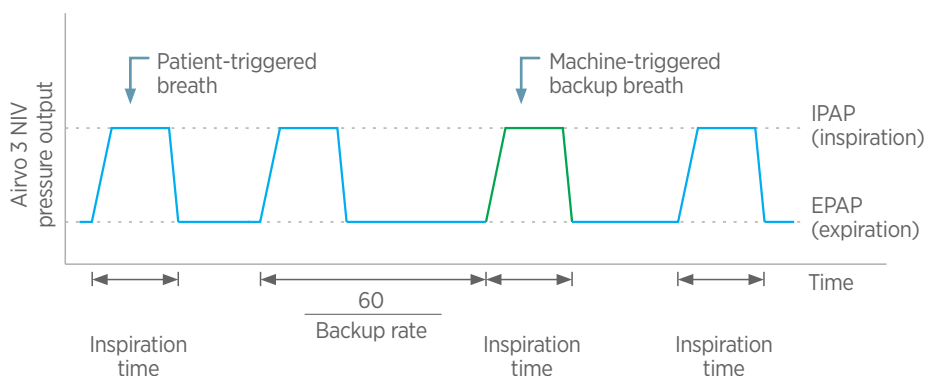
In Bi-level S/T mode, the duration of inhalation is controlled by the patient; the pressure drops from IPAP to EPAP when exhalation is detected. In Bi-level PCV mode, the duration of the breath is always equal to the Inspiration Time setting (Ti time). This difference is illustrated in the graphs above, which show the response of the Airvo 3 NIV to the same patient effort in Bi-level S/T and Bi-level PCV modes.



Bi-level S/T mode



Bi-level PCV mode

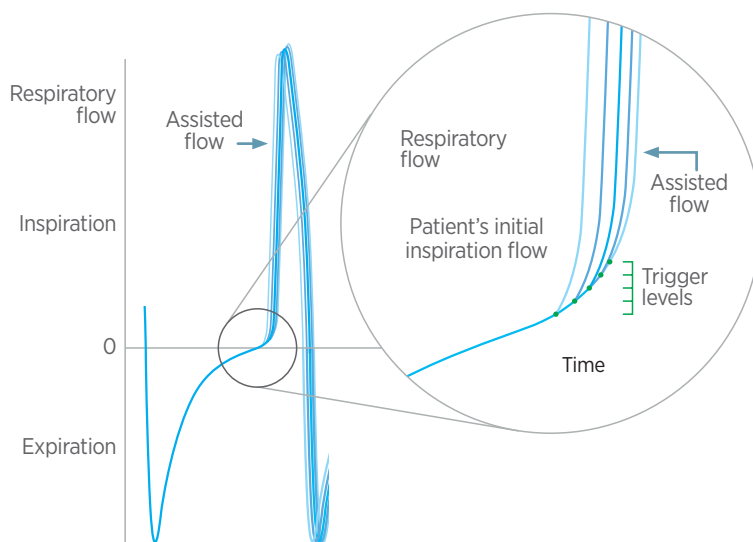


Machine-triggered backup breaths behave the same in both modes. Backup breaths occur whenever the patient's breathing rate drops below the Backup rate (see Backup breaths in section 11.1.4 above).

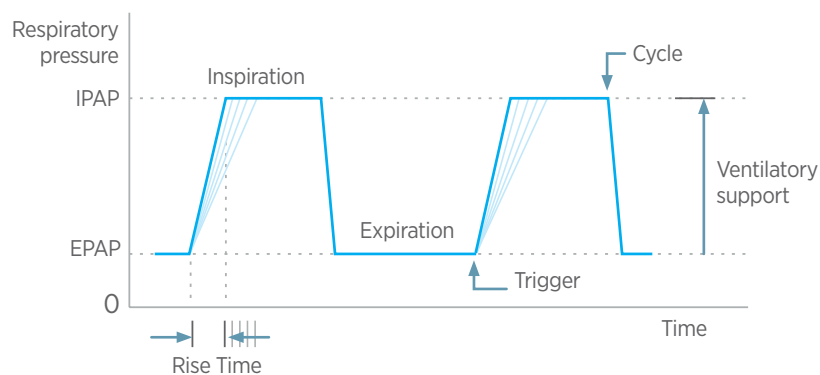
10.1.6 Triggering and cycling

Triggering is the detection of breaths spontaneously initiated by the patient, which causes the Airvo 3 NIV to begin increasing the respiratory pressure from the EPAP setting to the IPAP setting. The pressure changes at a rate controlled by the Rise Time. Respiratory pressure rises faster for lower values of Rise Time.

Cycling returns the respiratory pressure to the EPAP setting at the end of a breath. In Bi-level PCV mode, cycling occurs after the set Inspiration Time. In Bi-level S/T mode, the Airvo 3 NIV detects the change in the patient's respiratory flow as they start to breath out to cycle back to the EPAP setting.



Respiratory flow is assisted, by increasing respiratory pressure, when the patient's initial inspiration flow exceeds the trigger level.



Respiratory pressure shifts between EPAP and IPAP, providing pressure support to aid breathing.

Triggering and cycling both occur in the Bi-level S/T and Bi-level PCV modes; they do not apply to CPAP mode.

The Airvo 3 NIV uses the change in air-flow at the start of inspiration to trigger a breath. The sensitivity to changing air-flow can be adjusted to suit the patient's condition using the Trigger setting. This setting controls how much the air-flow must increase to trigger a breath:

- Decrease the trigger level if the Airvo 3 NIV is routinely missing breaths started by the patient.
- Increase the trigger level when the Airvo 3 NIV is routinely starting breaths that were not intended by the patient.

10.1.7 Leak compensation

A considerable volume of air leaks to the environment during noninvasive ventilation (NIV). There are two general sources of leaks:

1. Intentional leaks to vent CO₂ expired by the patient. The type of mask and respiratory pressure determine size of the intentional leak.
2. Unintentional leaks around the edges of the mask. Unintentional leaks change constantly with patient movement and breathing patterns.

The Airvo 3 NIV maintains respiratory pressure in the presence of leaks by adjusting the flow generator to compensate for both intentional and unintentional leaks.

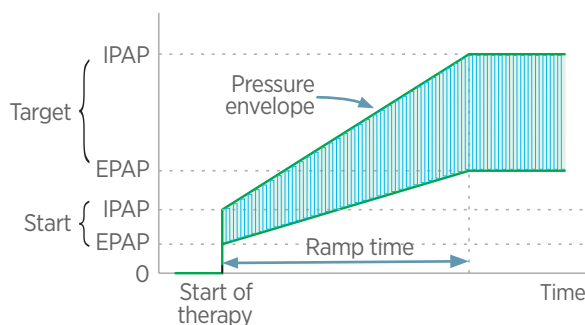
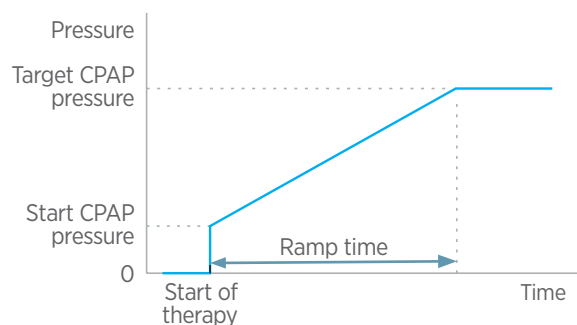
To maintain accurate triggering in the presence of unintentional leaks, the Airvo 3 NIV continuously measures the instantaneous pressure and flow rate of the respiratory gas delivered. By tracking these measurements across each breath, the leak flow and patient respiratory flow can be calculated and used to ensure accurate triggering of the next breath.

The leak rate is shown on the Data and Graphs  screen (see section 5.10). The unintentional leak rate is displayed when an F&P mask is selected and the total leak rate (intentional and unintentional combined) otherwise.

10.1.8 Ramping pressure

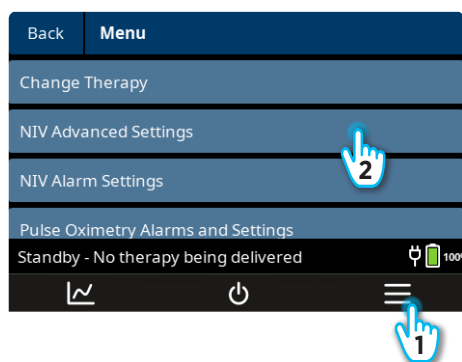
The Ramp setting may help your patients acclimatize to NIV therapy by reducing airway pressure temporarily. Airway pressure begins at a lower level when the ramp is started and increases slowly to the final pressure. The ramp function can be used in all three noninvasive ventilation modes. It behaves slightly differently in CPAP mode and the two Bi-level modes (Bi-level S/T and Bi-Level PCV):

- In CPAP mode, the respiratory gas is delivered at the selected Start Pressure when the ramp is started. The respiratory pressure is increased to the CPAP pressure over the duration set by the Ramp Time (5 – 45 minutes).
- In Bi-level S/T and Bi-level PCV modes, starting pressures for both EPAP and IPAP settings may be selected by the operator. Both EPAP and IPAP are separately increased from these initial values to the EPAP and IPAP target pressures respectively over the duration set by the Ramp Time (5 – 45 minutes).



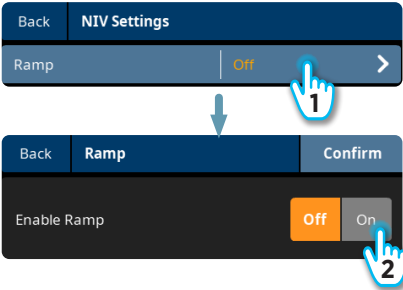
10.1.9 Using the pressure ramp

See ramping pressure section above for therapy behavior details during pressure ramps.



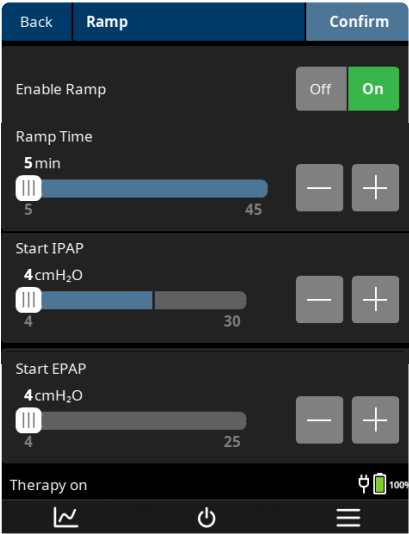
Open NIV settings

1. Tap  to open the system menu.
2. Select NIV Advanced Settings.



Enable the Ramp option

1. Tap Ramp to open the Ramp settings.
2. Select On to make the pressure ramp available.



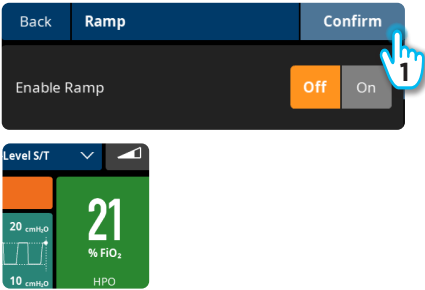
Adjust Ramp settings

Use the + / - buttons to select the required values for:

- Ramp Time: the duration of the ramp from the start to end pressure,
- Start IPAP: initial IPAP setting (Bi-level S/T and Bi-level PCV modes only),
- Start EPAP: initial EPAP setting (Bi-level S/T and Bi-level PCV modes only),
- Start CPAP: initial CPAP setting (CPAP mode only).

ⓘ Note

A Start IPAP/EPAP setting cannot be set higher than that final IPAP/EPAP setting.



Confirm Ramp settings

1. Tap Confirm to apply Ramp settings, or Back to discard any changes, and return to the NIV settings menu. Then tap Back twice to return to the Home screen.

The Ramp button is shown on the Home screen when ramp mode is enabled: tap the Ramp button to start pressure ramp and NIV therapy.

-  This shows the ramp is active. Tap the Ramp button to stop the pressure ramp and immediately apply the final target airway pressure settings.
-  This shows the ramp is off. Tap the Ramp button to start the pressure ramp.

Start NIV therapy without starting a pressure ramp by tapping the Start button. Stopping NIV therapy at any time stops the pressure ramp and resets the Ramp Time.

10.2 Alarm settings

The Airvo 3 NIV provides a set of alarms for NIV therapy. An alarm will be raised when measured values fall outside the user-selected limits.

Open NIV alarm settings by:

1. Tapping  to open the system menu,
2. Selecting NIV Alarm settings.

Individual NIV alarms can be disabled if they are not required for your patient. All alarm settings are persistent and will retain their previous value when the Airvo 3 NIV is turned on and same patient is selected. Selecting New Patient when reviewing the disinfection state applies the default values for your clinical environment to all alarms.

Name	Description	Default	Range	Unit
Low Respiratory Rate*	The average respiratory rate has dropped below the alarm threshold for more than 3 consecutive breaths.	Off	Off, 4 – 60	BPM
High Respiratory Rate*	The average respiratory rate has increased above the alarm threshold for more than 3 consecutive breaths.	Off	Off, 4 – 60	BPM
Low Minute Ventilation	The effective average minute ventilation has dropped below the alarm threshold for more than 3 breaths. Minute ventilation is the volume of air a patient moves in 1 minute.	Off	Off, 2 – 40	L/min
Low Inspiratory Pressure*	The delivered pressure during inhalation has fallen below the set inhalation pressure by the selected alarm threshold for more than 3 consecutive breaths.	Off	Off, 1 – 30	cmH ₂ O
High Inspiratory Pressure*	The delivered pressure during inhalation has risen above the set inhalation pressure by the selected alarm threshold for more than 3 consecutive breaths.	40	4 – 40	cmH ₂ O
Apnea**	A breath has not been triggered by either the patient or backup rate within the apnea alarm period. In Bi-level modes if a machine breath is triggered within the apnea alarm period the apnea timer is reset, preventing the apnea alarm from occurring.	Off	Off, 10 – 60	seconds

* The high alarm threshold cannot be set below the low alarm threshold.

Warning

Using different alarm settings on devices within a single area, such as an intensive care unit, can cause a hazard.

Caution

Setting alarm limits to extreme values can render the alarm system ineffective and may lead to patient injury.

Specifications

General

Dimensions	205 mm x 295 mm x 190 mm
Weight including battery	4.45 kg
Supply voltage/current	100 – 115 VAC, 2.4 A (2.6 A max¹) 220 – 240 VAC, 1.1 A (1.3 A max¹)
Supply frequency	50 – 60 Hz
USB port sourcing (1 and 2)	5 V, 0.25 A (maximum each port)
Auditory alarm	
Sound pressure level	> 40 dBA @ 1 m
Audio pause duration	120 seconds
Sound level	< 50 dBA @ 1 m
A-weighted sound power level	< 60 dBA
A-weighted sound pressure level	< 50 dBA
Ingress protection	IP22²
Expected service life	5 years³

Operating conditions

Ambient temperature	18 – 28 °C
Humidity	10 – 95% relative humidity (non-condensing)
Ambient pressure	700 – 1060 hPa
Altitude	
Optiflow:	0 – 3000m
NIV:	0 – 1000m
Mode of operation	Continuous operation
Maximum surface temperature of applied parts⁴	44 °C
Maximum delivered temperature of respiratory gas⁴	43 °C

Storage and transport conditions

Ambient temperature⁵,⁶	-10 – 50 °C
Humidity (non-condensing)	10 – 95% relative humidity

Battery (900PT957L)

Chemistry	Lithium Ion (Li-Ion)
Voltage	14.4 VDC
Capacity, Power output	99.4 Wh, 80 W
Battery life⁷	300 cycles or 2 years from first use (whichever comes first)
Recharge time	6 hours (maximum)
Shelf life	3 years
Operating time to 0%⁸	> 30 minutes
to 20%	
Typical	40 minutes
Worst case⁹	20 minutes (>30 minutes with flow and supplementary oxygen only)

Supplementary oxygen

Oxygen sensor startup time	< 30 s
Oxygen response time	< 60 s

Supplementary oxygen

Low-pressure oxygen (LPO) inlet port	
Line pressure	0 – 70 kPa
Maximum flow rate	60 L/min (STPD¹⁰)
% Concentration	93%, > 99%

Optiflow high flow therapy¹¹

Target humidity range	31 – 37 °C
Target flow range¹²	2 – 70 L/min
Maximum limited pressure¹³	60 cmH₂O
Maximum operating pressure	<45 cmH₂O
Oxygen concentration	21 – 100% FiO₂
Humidity⁴,⁷	> 33 mg/L at 37 °C target humidity, 10 – 60 L/min target flow¹⁴
Wall power	> 16 mg/L for all other settings
Static temperature stability	± 2 °C (All therapy modes)
Warm-up time¹⁵ (MR290 chamber)	
23 ± 2 °C to 37 °C	< 20 min

Noninvasive ventilation¹¹

Target humidity range	25 – 31 °C
Working pressure:	
CPAP range¹⁶	4 – 25 cmH₂O (+/- 1.5 cmH₂O + 5% of set pressure)¹⁷
IPAP range¹⁸	4 – 30 cmH₂O (+/- 1.5 cmH₂O + 5% of set pressure)¹⁷
EPAP range¹⁸	4 – 25 cmH₂O (+/- 1.5 cmH₂O + 5% of set pressure)¹⁷
Breath Rate	4 – 60 BPM (+/- 1 BPM)
Inspiration Time	0.3 to 4 s (+/- 0.1 s + 10% of set value)
Maximum limited pressure	60 cmH₂O
Maximum operating pressure	<60 cmH₂O
Intended delivery volume	> 50 mL
Humidity⁴,⁷	
Wall power	> 12 mg/L at 25-31 °C target humidity settings
Warm-up time¹⁵ (MR290 chamber)	
23 ± 2 °C to 37 °C	< 20 min

- The device is protected against solid objects larger than 12mm (e.g contact with a finger) and vertically dripping water will have no harmful effects when the enclosure is tilted at an angle of up to 15° from its standard position.
- Assumes typical usage pattern. Actual service life may vary.
- In accordance with ISO 80601-2-74. Tested to an accuracy of ± 1 °C or ± 1 mg/L, as appropriate.
- Storage at temperatures above 40 °C for prolonged periods will accelerate battery degradation.
- The device may require up to 24 hours to equilibrate to operating temperature before it is ready for use.
- For humidity performance under battery use, see Appendix 4.
- Worst-case battery time from fully charged to complete loss of power.
- Worst-case operating time applies to a fully charged battery at 25 °C that has experienced 3 years of storage followed by 300 charge/discharge cycles.
- Flow rate is expressed in STPD (standard temperature and pressure, dry) as per ISO 80601-2-74.
- Values are expressed in body temperature, pressure, saturated (BTPS), in accordance with ISO 80601-2-74, unless otherwise stated.
- Achievable flow rate depends on the patient interface selected.
- In accordance with ISO 80601-2-90.
- Applies to use with bypassed airway patient interfaces only.
- Applies when the device is connected to a wall power supply for warm-up.
- CPAP mode only.
- Dependent on a number of factors including mask fit and mask selection.
- Bi-level S/T and Bi-level PCV modes only.

1. Inrush current may reach 50 A.

Range and accuracy of measured parameters (Optiflow)

Measurement	Symbol	Displayed Range	Accuracy
Humidity	Temp	31 – 37 °C	Not specified
Flow rate	Flow	2 – 70 L/min	± (1 + 5% of reading) L/min*
Oxygen concentration*	FiO ₂	21 – 100%	Lower of: ± 4%, or ± (2.5% + 2.5% of reading) - excluding rounding to 21% and 100%, as appropriate - provided “Oxygen concentration” setting is correct
Respiratory rate	RR	4 – 70 BPM	RMS error of 3 BPM**
Peripheral blood oxygen saturation	SpO ₂	1 – 100%	See Nonin pulse oximetry specifications below.
Pulse rate	PR / 	Masimo 25-240 beats/min Nellcor 20-300 beats/min Nonin 18–321 beats/min	See Nonin pulse oximetry specifications below.
Perfusion Index	PI	0.02%-20%	Not specified (Masimo only)

* Oxygen measurements are automatically compensated for changes in barometric pressure.

** An RMS accuracy is a statistical calculation of the difference between device measurements and reference measurements. Approximately two-thirds of the device measurements fell within +/- ARMS of the reference measurements in a controlled study.

Range and accuracy of measured parameters (NIV)

Measurement	Symbol	Displayed Range	Accuracy
Patient Leak	Pt. Leak	0 – 70 L/min BTPS	± 15 L/min
Total Leak	TTL. Leak	0 – 100 L/min BTPS	Higher of: ± 15 L/min or ± 20% of reading, at 10 cmH ₂ O
Oxygen concentration* **	FiO ₂	21 – 100%	Lower of: ± 4%, or ± (2.5% + 2.5% of reading) - excluding rounding to 21% and 100%, as appropriate - provided “Oxygen concentration” setting is correct
Respiratory rate	RR	4 – 60 BPM	± 1 BPM
Exhaled tidal volume	Vt	200 – 2000 ml BTPS	20% of reading
Minute ventilation	MV	2 – 40 L/min	Higher of: ± 1 L/min or 20% of reading
% Trigger breaths	Pt. Trig	0 – 100%	± 10%
T _i /T _{tot}	Ti/Ttot	10 – 80%	± 10%
Flow waveform	N/A	-50 – 150 L/min	Higher of: ± 15 L/min or 20% of reading
Pressure waveform	N/A	0 – 40 cmH ₂ O	± (1.5 cmH ₂ O + 5% of set pressure)
Volume waveform	N/A	200 – 2000 ml BTPS	± 20% of actual reading

* Test equipment and methods are selected and controlled to ensure the uncertainty coverage is no more than 30% of the disclosed tolerance.

** Oxygen measurements are automatically compensated for changes in barometric pressure.

Pulse oximetry specifications (Masimo)

Specifications are tabulated for the Airvo 3 NIV and all compatible sensors unless otherwise stated.

Masimo:

Data update period	< 30 sec
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Measurement wavelengths and Output Power	Radiant power with a 50mA pulsed LED is less than 15mW. Masimo's RD SET and LNCS sensors use red and infrared light emitting diodes. The wavelengths for all sensors except TC-I and TF-I sensors are 660 Nanometres (nm), and 905nm for red and infrared respectively. TC-I: 653nm and 880nm for red and infrared respectively. TF-I: 660nm and 880nm for red and infrared respectively. This information is especially useful for clinicians performing photodynamic therapy.
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Accuracy (see notes 1-12 below)

Saturation (%SpO₂) - During No Motion Conditions

Adults/Pediatrics	70-100% ± 2 digits 0-69% unspecified
Neonates	70-100% ± 3 digits 0-69% unspecified

Saturation (%SpO₂) - During Motion Conditions

Adults/Pediatrics	70-100% ± 3 digits 0-69% unspecified
Neonates	70-100% ± 3 digits 0-69% unspecified

Pulse Rate (bpm) - During No Motion Conditions

Adults, Pediatric, Neonates	25 to 240 ± 3 digits
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Pulse Rate (bpm) - During Motion Conditions

Adults, Pediatric, Neonates	25 to 240 ± 5 digits
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Resolution

Saturation (%SpO₂)	1%
Pulse Rate (bpm)	1

Low Perfusion Performance

Pulse Amplitude	± 2 digits
% Transmission	5%
Saturation (%SpO₂)	± 2 digits
Pulse rate	± 3 digits

- The Masimo SET technology with Masimo sensors has been validated for no motion accuracy in human blood studies on healthy adult male and female volunteers with light to dark skin pigmentation in induced hypoxia studies in the range of 70-100% SpO₂ against a laboratory CO-Oximeter and ECG monitor. This variation equals ±1 standard deviation. Plus or minus one standard deviation encompasses 68% of the population.
- The Masimo SET technology with Masimo sensors has been validated for motion accuracy in human blood studies on healthy adult male and female volunteers with light to dark skin pigmentation in induced hypoxia studies while performing rubbing and tapping motions, at 2 to 4 Hz at an amplitude of 1 to 2 cm and a non-repetitive motion between 1 to 5 Hz at an amplitude of 2 to 3 cm in induced hypoxia studies in the range of 70-100% SpO₂ against a laboratory CO-Oximeter and ECG monitor. This variation equals ±1 standard deviation, which encompasses 68% of the population.
- The Masimo SET technology has been validated for low perfusion accuracy in bench top testing against a Bioteck Index 2™ simulator and Masimo's simulator with signal strengths of greater than 0.02% and transmission of greater than 5% for saturations ranging from 70 to 100%. This variation equals ±1 standard deviation. Plus or minus one standard deviation encompasses 68% of the population.
- The Masimo SET Technology with Masimo Neo sensors has been validated for neonatal motion accuracy in human blood studies on healthy adult male and female volunteers with light to dark skin pigmentation in induced hypoxia studies while performing rubbing and tapping motions, at 2 to 4 Hz at an amplitude of 1 to 2 cm and a non-repetitive motion between 1 to 5 Hz at an amplitude of 2 to 3 cm in induced hypoxia studies in the range of 70-100% SpO₂ against a laboratory CO-Oximeter and ECG monitor. This variation equals ±1 standard deviation, which encompasses 68% of the population. 1% has been added to the results to account for the effects of fetal hemoglobin present in neonates.
- The Masimo SET technology with Masimo sensors has been validated for pulse rate accuracy for the range of 25 -240 bpm in bench top testing against a Bioteck Index 2™ simulator. This variation equals ±1 standard deviation. Plus or minus one standard deviation encompasses 68% of the population.
- See sensor directions for use (DFU) for complete application information. Unless otherwise indicated, reposition reusable sensors at least every 4 hours and adhesive sensors at least every 8 hours.
- The TC-I sensor is contraindicated for patients with pierced ears at the measuring site.
- The TF-I sensor must be removed and repositioned to a different monitoring site at least every 2 hours. If extended monitoring is required, use of a single patient adhesive digit sensor is recommended.
- The TF-I, TC-I, and DBI sensors were not validated under motion conditions.
- The Trauma and Newborn sensors are for use only with instruments containing Masimo SET oximetry (Version 4.1.0.1 or higher) or monitors licensed to use specialty sensors.
- Sensor accuracy specified when used with Masimo technology using a Masimo patient cable for LNOP sensors, RD SET sensors, the LNCS sensors, or the M-LNCS sensors. Numbers represent Arms (RMS error compared to the reference). Because pulse oximeter measurements are statistically distributed, only about two-thirds of the measurements can be expected to fall within a range of ± Arms compared to the reference value. Unless otherwise noted, SpO₂ accuracy is specified from 70% to 100%. Pulse Rate accuracy is specified from 25 to 240 bpm.
- Masimo M-LNCS, LNOP, RD, and LNCS sensors, cables, and adapters have the same optical and electrical properties and may differ only in application type (adhesive/non-adhesive/hook & loop), cable lengths, optical component locations (top or bottom of sensor as aligned with cable), adhesive material type/size, and connector type (LNOP 8 pin modular plug, RD 15 pin modular plug, LNCS 9 pin, cable based, and M-LNCS 15 pin, cable based). All sensor accuracy information and sensor application instructions are provided with the

associated sensor directions for use.

- 13 For M-LNCS Blue, LNOP Blue is the predicate. Masimo SET technology with LNOP Blue sensors have been validated for no motion accuracy in human blood studies on neonatal, infant and pediatric patients with congenital cyanotic cardiac lesions in the range of 60%-100% SpO₂ against a laboratory co-oximeter. This variation equals plus or minus one standard deviation, which encompasses 68% of the population.
- 14 The presented 510(k) reference is based on the specific FDA clearance for the specific Masimo technology board cleared with the compatible Masimo sensor. The 510(k) reference may vary for the Masimo sensor depending on the pulse oximetry technology (i.e. Masimo SET, Masimo rainbow SET, Philips FAST, Nellcor).

Pulse oximetry specifications (Nellcor)

Please refer to the Oxicable Instructions for Use for product specifications

Pulse oximetry specifications (Nonin)

Specifications are tabulated for the Airvo 3 NIV and all compatible sensors unless otherwise stated.

Nonin:

Data update period	< 30 sec	
Measurement wavelengths and Output Power*	Red: 660 nanometers @ 0.8 mW max. avg. Infrared: 910 nanometers @ 1.2 mW max avg. (using Nonin Purelight® sensor)	
SpO₂ Accuracy (A_{rms}**)	70 to 100%	
No Motion	Adults/Pediatrics***	Neonates
Reusable		
8000AX Series:	± 2 digits	N/A
800XJ Series:	± 3 digits	N/A
8000SX Series:	± 2 digits	N/A
8000R:	± 3 digits	N/A
8000Q2:	± 3 digits	N/A
Disposable		
6000CX Series:	± 2 digits	± 3 digits
7000X Series:	± 2 digits	± 3 digits
Motion		
Reusable		
8000AX Series:	± 2 digits	N/A
800XJ Series:	± 3 digits	N/A
8000SX Series:	± 3 digits	N/A
Low Perfusion****	± 2 digits	± 3 digits
Pulse Rate Accuracy	Adults/Pediatrics***	Neonates
No Motion (18 - 300 BPM)		
Reusable		
8000AX Series:	± 3 digits	N/A
800XJ Series:	± 3 digits	N/A
8000SX Series:	± 3 digits	N/A
8000R:	± 3 digits	N/A
8000Q2:	± 3 digits	N/A
Disposable		
6000CX Series:	± 3 digits	± 3 digits
7000X Series:	± 3 digits	± 3 digits
Motion (40 - 240 BPM)		
Reusable		
8000AX Series:	± 5 digits	N/A
800XJ Series:	± 5 digits	N/A
8000SX Series:	± 5 digits	N/A
Low Perfusion (40 - 240 BPM)****	± 3 digits	± 3 digits

* This information is especially useful for clinicians performing photodynamic therapy.

** ± 1 A_{rms} represents approximately 68% of measurements.

*** Includes Infant patients

**** Does not apply to those sensors listed as N/A under the neonate column, 8000R and 8000Q2

Notes:

- SpO₂ accuracy testing is conducted during induced hypoxia studies on healthy, non-smoking, light-to-dark-skinned subjects during motion and no-motion conditions in an independent research laboratory. The measured arterial hemoglobin saturation value (SpO₂) of the sensors is compared to arterial hemoglobin oxygen (SaO₂) value, determined from blood samples with a laboratory co-oximeter. The accuracy of the sensors in comparison to the co-oximeter samples measured over the SpO₂ range of 70 – 100%. Accuracy data is calculated using the root-mean-squared (Arms value) for all subjects, per ISO 80601-2-61 formerly ISO 9919, Standard Specification for Pulse Oximeters for Accuracy.
- Pulse rate motion testing measures pulse rate accuracy with motion artifact simulation introduced by a pulse oximeter tester. This test determines whether the oximeter meets the criteria of ISO 80601-2-61, formerly ISO 9919, for pulse rate during simulated movement, tremor, and spike motions.
- Low perfusion testing uses an SpO₂ Simulator to provide a simulated pulse rate, with adjustable amplitude settings at various SpO₂ levels. The module must maintain accuracy in accordance with ISO 80601-2-61, formerly ISO 9919, pulse rate and SpO₂ at the lowest obtainable pulse amplitude (0.3% modulation).
- The Nonin Xpod has been validated for pulse rate accuracy from 18–300 bpm with no motion and from 40–240 bpm with motion. Testing was carried out using a Datrend Oxitest Plus 7 simulator.
- An SpO₂ Simulator was used to provide a simulated pulse rate, with adjustable amplitude settings at various SpO₂ levels. The module must maintain accuracy in accordance with ISO 80601-2-61, formerly ISO 9919, pulse rate and SpO₂ at the lowest obtainable pulse amplitude (0.3% modulation).

Standards compliance

Designed to conform to the following standards:

IEC 60601-1:2005+AMD1:2012 +AMD2:2020

IEC 60601-1-2:2014+AMD1:2020



Medical — cardio, vascular and pulmonary equipment as to electrical shock, fire and mechanical hazards only in accordance with AAMI ES60601-1 (2005) + AMD 1 (2012), CSA CAN/CSA-C22.2 NO. 60601-1:14, IEC 60601-1-6:2010, AMD1:2013, IEC 60601-1-8: 2006 + Am.1: 2012, ISO 80601-2- 61:2017, COR1:2018, ISO 80601-2-74:2017

ANSI/AAMI ES 60601-1:2005 and A1:2012 and A2:2021

CAN/CSA-C22.2 NO.60601-1:14+A2:2022 (R2022)

IEC 60601-1-6:2010+AMD1:2013+AMD2:2020

IEC 60601-1-8:2006+AMD1:2012+AMD2:2020

IEC 60601-1-10:2007+AMD1:2013+AMD2:2020

ISO 80601-2-61:2017

ISO 80601-2-74:2021

Do not place any part of the device or accessories within 30 cm of any portable mobile radio frequency communication equipment. The Airvo 3 NIV complies with the electromagnetic compatibility requirements of IEC 60601-1-2. In certain circumstances the Airvo 3 NIV may affect or be affected by nearby equipment because of electromagnetic interference. Excessive electromagnetic interference may affect the therapy delivered by the device. If this should happen, try moving the Airvo 3 NIV or the unit causing interference, or consult your healthcare provider.

FCC compliance

This device has been tested and found to comply with the limits for a Class B digital device, pursuant to part 15 of the FCC Rules. These limits are designed to provide reasonable protection against harmful interference in a residential installation. This device generates, uses, and can radiate radio frequency energy and, if not installed and used in accordance with the instructions, may cause harmful interference to radio communications. However, there is no guarantee that interference will not occur in a particular installation. If this device does cause harmful interference to radio or television reception, which can be determined by turning the device off and on, the user is encouraged to try to correct the interference by one or more of the following measures:

- Reposition or relocate the receiving antenna.
- Increase the separation between the device and receiver.
- Connect the device into an outlet on a circuit different from that to which the receiver is connected.
- Consult your healthcare provider or your Fisher & Paykel Healthcare representative for help.

Accessory equipment connected to the any port of the Airvo 3 NIV must be certified to IEC 60601-1-1 or IEC 60950-1.

All configurations shall comply with the system standard IEC 60601-1-1. Anyone who connects additional equipment to the signal input part or signal output part configures a medical system and is therefore responsible for ensuring that the system complies with the requirements of the system standard IEC 60601-1-1. If in doubt, consult your technical services department or your local Fisher & Paykel Healthcare representative. Certain elements of the software included with product are supplied under the license terms of third parties, including elements of the software that are subject to certain open source software licenses. Where required by the terms of these licenses, Fisher & Paykel Healthcare Limited provides notices regarding such software elements on its website. Please visit www.fphcare.com/airvo3/third-party-licenses to view these notices. Note that the notices that apply may be updated as the software included in the product is updated. The F&P Airvo 3 NIV is in compliance with Directive 2014/53/EU. The full text of the EU declaration of conformity is available at the following internet address: www.fphcare.com/certifications.

Device disposal instructions



This device contains electronic components and a lithium-ion battery. Regulations in your country may require them to be collected for waste recovery and recycling to reduce the environmental impact. Please dispose of this device in accordance with local regulations.

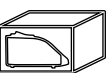
Disposal of accessories, spare parts and packaging



Dispose of accessories, spare parts and packaging according to local guidelines. Place the breathing tube and water chamber in a waste bag at the end of use and discard with normal waste. Hospitals should discard according to their standard method for disposing of contaminated product.

Glossary

Symbols

 For safety reasons, refer to the instructions for use	 Warning, hot surface	 Power on/off button	 System menu button
 Alarm symbol	 Alarm limits	 USB port and Compatible USB device detected	IP22 Protected against ingress of small objects and water drops
 Class II equipment (double insulated)	 Magnetic Resonance (MR) unsafe	 Non-ionizing electromagnetic radiation	 Humidity range
 Do not use if package is damaged	 Type BF applied part (body floating)	 Do not discard as regular waste	 Temperature range
 Operating conditions	 Storage and transport conditions	 Importer	 Distributor
 Catalogue number	 YYYY-MM-DD Date of manufacture	 Manufacturer	 YYYY-MM-DD Manufacturer and date of manufacture
 Serial number	 Medical Device*	CE 0123 European conformity - TÜV SÜD*	 Regulatory compliance mark*
 UL Classified Mark Canada, USA*	 EU Authorised representative	 Contains hazardous substances	

*symbol seen on select models

Appendix 1. Patient consumables

The patient interfaces and accessories shown in the tables below are approved for use with the Airvo 3 NIV. Carefully read the user instructions, including all warnings and cautions, supplied with each device before use.

Some accessories may not be available in certain countries. Contact your Fisher & Paykel Healthcare representative for the latest information on patient interfaces available for the Airvo 3 NIV. All patient interfaces are Type BF applied parts.

Optiflow high flow therapy

Description	Part number	Size	Pack size
Optiflow+ nasal interface	OPT942	Small	20
	OPT944	Medium	20
	OPT946	Large	20
Optiflow+ Duet interface	OPT962	Small	20
	OPT964	Medium	20
	OPT966	Large	20
Optiflow 3S nasal interface	OPT1042	Small	20
	OPT1044	Medium	20
	OPT1046	Large	20
Optiflow Junior 2 nasal interface*	OJR414 (WJR112)	Medium	20 (20)
	OJR416 (WJR112)	Large	20 (20)
	OJR418 (WJR112)	Extra Large	20 (20)
Optiflow Junior 2+ nasal interface*	OJR520 (WJR114)	XXL	10 (10)
Optiflow Junior 2 WigglewiNG	WJR212	M, L, XL	20
	WJR214	XXL	10
Optiflow+ tracheostomy direct connection	OPT970	15 mm	20
Optiflow+ mask interface adapter**	OPT980	22 mm mask interface adapter	20
AirSpiral tube and chamber kit	900PT561	—	10
AirvoNeb tube and chamber kit	900PT562	—	10

* Wigglepads part numbers are shown in parentheses.

** The mask adapter interface is designed for vented masks only. Do not use sealed masks with Optiflow high flow therapy

* An exhalation port must be included in the breathing circuit when using non-vented NIV patient interfaces. An exhalation port is included in the AirSpiral NIV tube and chamber kit.

Adult NIV therapy

Some accessories may not be available in certain countries. Please contact your local Fisher & Paykel Healthcare representative for more information.

Description	Part number	Size	Pack size
AirSpiral NIV tube and chamber kit	900PT573	—	10
NIV breathing tube filter	900PT510	—	20
F&P FreeMotion™ vented Hospital Full Face mask	RT040S	Small	10
	RT040M	Medium	10
	RT040L	Large	10
F&P FreeMotion vented Hospital Nasal mask	RT042	Medium	10
	RT042L	Large	10
F&P FreeMotion non-vented Hospital Full Face mask*	RT043S	Small	10
	RT043M	Medium	10
	RT043L	Large	10
F&P Nivairo™ Non-vented Hospital Full Face Mask, Compatible with Single-limb Circuits, Requiring Exhalation Port	RT045XS	Extra Small	10
	RT045S	Small	10
	RT045M	Medium	10
	RT045L	Large	10
F&P Nivairo Vented Hospital Full Face Mask, Compatible with Single-limb Circuits	RT047XS	Extra Small	10
	RT047S	Small	10
	RT047M	Medium	10
	RT047L	Large	10
F&P Nivairo+™ Non-vented Hospital Full Face Mask, Compatible with Single-limb Circuits, Requiring Exhalation Port	RT055XS	Extra Small	10
	RT055S	Small	10
	RT055M	Medium	10
	RT055L	Large	10
F&P Nivairo+ Vented Hospital Full Face Mask, Compatible with Single-limb Circuits	RT057XS	Extra Small	10
	RT057S	Small	10
	RT057M	Medium	10
	RT057L	Large	10

* An exhalation port must be included in the breathing circuit when using non-vented NIV patient interfaces. An exhalation port is included in the AirSpiral NIV tube and chamber kit.

Adult NIV therapy

Some accessories may not be available in certain countries. Please contact your local Fisher & Paykel Healthcare representative for more information.

Description	Part number	Size	Pack size
F&P Visairo™ Non-vented Hospital Under Nose Mask, compatible with Single-limb Circuits, Requiring Exhalation Port	RT075A	A	10
	RT075B	B	10
	RT075C	C	10
F&P Visairo Vented Hospital Under Nose Mask, Compatible with Single-limb Circuits	RT077A	A	10
	RT077B	B	10
	RT077C	C	10
F&P OptiNIV™ Hospital Vented Mask Compatible with Single-limb Circuits	ONIV117A	A	10
	ONIV117B	B	10
	ONIV117C	C	10
F&P Simplus™ Mask - Small	400475	Small	1
F&P Simplus Mask - Medium	400476	Medium	1
F&P Simplus Mask - Large	400477	Large	1

Appendix 2. Parts and accessories

Some accessories may not be available in certain countries. Please contact your local Fisher & Paykel Healthcare representative for more information.

Accessories

Description	Part number
Mobile pole stand	900PT421
Mobile pole stand handle	900PT445
Mobile pole stand clamp	900PT428
Oxygen-bottle holder	900PT427, 900PT427L
Storage basket	900PT426
Storage cover	900PT603
HPO Dual-Input Manifold (DISS, NIST, SIS)	900PT460D, 900PT460N, 900PT460S
HPO adapter (DISS to NIST)	900PT462DN
Airvo 3 data port adapter*	900PT473
Airvo 3 USB service cable	900PT474
Fisher & Paykel Healthcare Device Manager	900PT475
Disinfection kit**	900PT600

* The data port adapter allows for data to be transferred from the Airvo 3 to patient monitoring and hospital computer systems. Integration is required to enable functionality.

**A disinfection kit is required when using the built-in disinfection mode to disinfect the outlet elbow. It is not required for hospitals using a washer-disinfector to clean and disinfect the outlet elbow.

Warnings

Equipment connected to Airvo 3 via the Data Port Adaptor (900PT473) must be certified according to IEC 60950-1 or IEC 60601-1. All combinations of equipment must be in compliance with IEC 60601-1 requirements for medical electrical systems. Any person who connects additional equipment has now configured a medical system and is therefore responsible for ensuring that the system complies with the requirements of IEC 60601-1. Consult a technical specialist in your hospital for more information.

Using accessories other than those specified may result in increased electromagnetic emissions or decreased electromagnetic immunity of the Airvo 3.

All data transferred from Airvo 3 is intended for information only and must not be used as the sole basis for diagnostic or therapeutic decisions.

Spare parts

Description	Part number
Cleaning sponge sticks	900PT602
Storage cover	900PT603
Outlet elbow	900PT930
Air Filter	900PT933
Battery module	900PT957L

Appendix 3. Pulse oximetry accessories

The pulse oximetry accessories listed below are compatible with the Airvo 3. Carefully read the user instructions, including all warnings and cautions, supplied with each device before use. Not all accessories are available in all markets, and some accessories may not be available from Fisher & Paykel Healthcare.

Masimo:

Part numbers of compatible Masimo pulse oximetry USB connector cable, adapters, and extension cables

Description	Masimo part number (cable length)
uSpO ₂ Masimo SET Oximetry Cable, USB	3412 (1.8m)
RD to LNC Adapter Cable	4089 (0.9m)
RD to LNC Adapter Cable	4105 (0.45m)
LNC-4-Ext	2021 (1.2m)

Part numbers of compatible Masimo pulse oximetry sensor cables and sensor consumables

Sensor description	Masimo part number (cable length) (other information)
RD SET™ DCI Series Adult Reusable Finger Clip Sensors	4050 (0.9m)
RD SET™ DCI-P Series Pediatric Reusable Finger Clip Sensors	4051 (0.9m)
RD SET™ TC-I Reusable Tip Clip Sensor	4053 (0.9m)
RD SET™ YI SpO ₂ Multisite Reusable Sensor	4054 (0.9m)
RD SET™ TF-I SpO ₂ Reusable Transflectance Forehead Sensor	4055 (0.9m)
RD SET™ DB-I Reusable Soft Sensors	4052 (0.9m)
RD SET™ Series Adt SpO ₂ Disposable Sensors	4000 (0.45m) (20 pack)
RD SET™ Series Pdt SpO ₂ Disposable Sensors	4001 (0.45m) (20 pack)
RD SET™ Series Inf SpO ₂ Disposable Sensors	4002 (0.45m) (20 pack)
RD SET™ Series Neo SpO ₂ Disposable Sensors	4003 (0.45m) (20 pack)
RD SET™ Series Neo Pt SpO ₂ Disposable Sensors	4004 (0.45m) (20 pack)
RD SET Series Neo Pt SpO ₂ Disposable Sensors (Non-Adhesive)	4005 (0.45m) (20 pack)
RD SET™ Speciality Sensor Series Adult Trauma	4011 (10 pack)
RD SET™ Specialty Sensor Series Newborn Neonatal	4013 (10 pack)
RD SET™ Specialty Sensor Series Newborn, Infant, Pediatric	4012 (10 pack)
RD SET™ Blue Disposable Sensor	4014 (10 pack)
RD SET™ Ear Sensor	4015 (0.9m) (10 pack)
RD SET™ TFA-I SpO ₂ Disposable Transflectance Forehead Sensor	4016 (0.9m)
LNCS® DCI ADT Reusable Sensor	1863 (0.9m)
LNCS® DCIP Reusable Sensor	1864 (0.9m)
LNCS® TC-I Reusable Tip Clip Sensor	1895 (0.9m)

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Masimo:**Part numbers of compatible Masimo pulse oximetry sensor cables and sensor consumables**

Sensor description	Masimo part number (cable length)
LNCS[®] YI SpO₂ Multisite Reusable Sensor	2258 (0.9m)
LNCS[®] TF-I[®] Adult SpO₂ Reusable Transflectance Sensor	1896 (0.9m)
LNCS[®] DB-I Series Reusable Soft Sensors	2653 (0.9m)
LNCS[®] Adtx, Adult Adhesive Sensor	1859 (0.45m) (20 pack)
LNCS[®] Adtx-3, Adult Adhesive Sensor	2317 (0.9m) (20 pack)
LNCS[®] Pdtx, Pediatric Adhesive Sensor	1860 (0.45m) (20 pack)
LNCS[®] Pdtx-3, Pediatric Adhesive Sensor	2318 (0.9m) (20 pack)
LNCS[®] Inf, Infant Adhesive Sensor	2328 (0.45m) (20 pack)
LNCS[®] Inf-3, Infant Adhesive Sensor	2319 (0.9m) (20 pack)
LNCS[®] Inf-L, Infant Adhesive Sensor	1861 (0.9m) (20 pack)
LNCS[®] Neo, Neonate Adhesive Sensor	2329 (0.45m) (20 pack)
LNCS[®] Neo-3, Neonate Adhesive Sensor	2320 (0.9m) (20 pack)
LNCS[®] Neo-L, Neonate Adhesive Sensor	1862 (0.9m) (20 pack)
LNCS[®] NeoPt, Sensitive Skin Neonate Adhesive Sensor	2330 (0.45m) (20 pack)
LNCS[®] NeoPt-3, Sensitive Skin Neonate Adhesive Sensor	2321 (0.9m) (20 pack)
LNCS[®] NeoPT-L, Sensitive Skin Neonate Adhesive Sensor	1901 (0.9m) (20 pack)
LNCS[®] NeoPt-500, Neonate Non-Adhesive Sensor	2331 (0.45m) (20 pack)
LNCS[®] Trauma Adult Adhesive Sensor	2411 (0.9m) (20 pack)
LNCS[®] Specialty Sensor Series Newborn neonatal	2412 (0.9m) (20 pack)
LNCS[®] Specialty Sensor Series Newborn, Infant, Pediatric	2413 (0.9m) (20 pack)
LNCS[®] E1 Ear Sensor	2918 (0.9m) (10 pack)
LNCS[®] TFA-1 SpO₂ Disposable Transflectance Forehead Sensor	3858 (0.9m) (10 pack)

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Nellcor:**Adaptors and extension cables**

Description	Part number
Pulse Oximetry Connection Converter 1	900PT476 (0.3m)
Medtronic/Nellcor Oxicable	PMC10N-SF (2.8m)

Part numbers of compatible Nellcor pulse oximetry sensor consumables

Sensor description	Nellcor part number (cable length) (other information)
Nellcor SpO₂ Forehead Sensor	MAXFAST (0.75m) (24 pack)
Nellcor SpO₂ Nonadhesive Sensor	SC-A/SC-NEO/SC-PR (0.9m) (24 pack)
Nellcor Flexible SpO₂ Sensor	FLEXMAX/FLEXMAX-P (0.9m)
Nellcor SpO₂ Adhesive Sensors	MAXA/MAXAL/MAXN/MAXI/MAXP (0.9m) (24 pack)
Nellcor SpO₂ Adhesive Sensors Nasal	MAXR (0.45m) (24 pack)
Nellcor Reusable SpO₂ Sensors	DS100A (1 pack), OXI-A, OXI-N, OXI-P, OXI-I (24 pack) (0.9m)
Nellcor Reusable Multisite SpO₂ Sensors	D-YS, D-YSE, D-YSPD (0.9m), PDSLVS (replacement sleeves, 12 pack)
Nellcor Single-Patient Use Sensor Wraps	POSEY (for OXI-A/N/P/I, D-YS) (12 pack)
Nellcor Single-Patient Use Adhesive Sensor Wraps	ADH-A/N (for OXI-A/N, D-YS), ADH-P/I (for OXI-P/I, D-YS) (100 pack)
Nellcor Single-Patient Use Foam Sensor Wraps	FOAM A/N (for OXI-A/N, D-YS), FOAM P/I (for OXI-P/I, D-YS) (100 pack)

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Nonin:**Part numbers of compatible Nonin pulse oximetry USB connector cables**

Description	Nonin part number (cable length)
Nonin Xpod 3012HR USB Connector Cable	114403-001 (1m)
Xpod® 3012 LP with USB Connector	6703-001 (1m)

Part numbers of compatible Nonin pulse oximetry sensor cables and sensor consumables

Sensor description	Nonin part number (cable length) (pack size)
8000SS reusable soft sensors, small	6837-000 (1m), 6837-300 (3m)
8000SM reusable soft sensors, medium	6836-000 (1m), 6836-300 (3m)
8000SL reusable soft sensors, large	6835-000 (1m), 6835-300 (3m)
8000AA adult reusable finger clip sensors	3278-001 (1m), 3278-006 (2m, 6.6m), 3278-003 (3m)
8000AP pediatric reusable finger clip sensors	2360-000 (1m), 2360-003 (3m)
8000Q2 ear clip sensor	6455-000 (1m)
8000R reflectance sensor	0487-000 (1m)
8000J adult semi-reusable Flex Sensor	0741-000 (1m), 2353-002 (3m) (includes x25 8000JFW FlexiWraps®)
8008J infant semi-reusable Flex Sensor	0740-000 (1m) (includes x25 8008JFW FlexiWraps)
8001J neonatal semi-reusable Flex Sensor	0739-000 (1m) (includes x25 8001JFW FlexiWraps)
6000CA adult cloth disposable sensors	7426-001 (1m) (24 pack)
6000CP pediatric cloth disposable sensors	7426-002 (1m) (24 pack)
6000CI infant cloth disposable sensors	7426-003 (1m) (24 pack)
6000CN neonatal cloth disposable sensors	7426-004 (1m) (24 pack)
7000A adult Flexi-Form® III disposable sensors	7427-001 (1m) (24 pack)
7000P pediatric Flexi-Form III disposable sensors	7427-002 (1m) (24 pack)
7000I infant Flexi-Form III disposable sensors	7427-003 (1m) (24 pack)
7000N neonatal Flexi-Form III disposable sensors	7427-004 (1m) (24 pack)
8000JFW adult FlexiWraps	4097-000, (25 pack), for use with 8000J
8008JFW infant FlexiWraps	4774-000, (25 pack), for use with 8008J
8001JFW neonatal FlexiWraps	4777-000, (25 pack), for use with 8001J
8000H reflectance sensor holder pack	0616-000, (10 caps & 20 adhesive stickers) for use with 8000R
Sensor Clip for LP Xpod External Pulse Oximeter	7504-001

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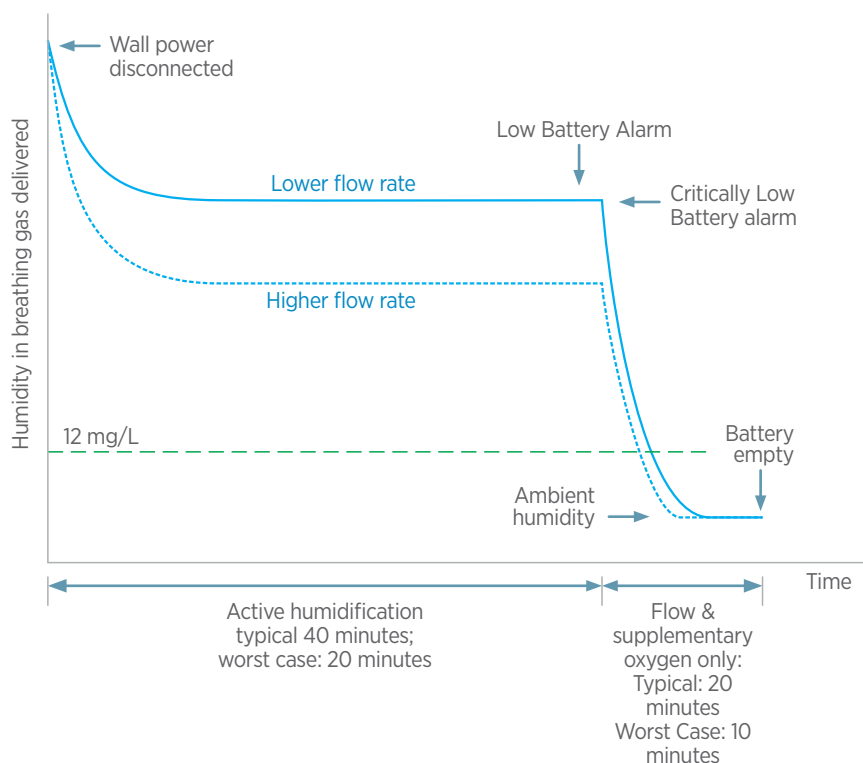
Appendix 4. Humidification behavior during battery operation

The Airvo 3 NIV reduces the energy used to humidify breathing gases when not powered from a wall power supply, to conserve battery power. In all cases, the Airvo 3 NIV continues supplying supplementary oxygen and breathing gases until the battery is depleted.

For Optiflow high flow therapy, active humidification of the breathing gases is reduced during battery operation. If the Critically Low Battery alarm is raised, active humidification is stopped to conserve battery power.

For NIV therapy, active humidification of the breathing gases is stopped during all battery operation to conserve battery power.

Connect the Airvo 3 NIV to a wall power supply before the battery is empty to automatically resume normal therapy. If the Airvo 3 NIV battery is depleted, the device stops supplying supplementary oxygen and breathing gases, powers down and produces the Power Out alarm. To resume therapy after the device has powered down, connect the Airvo 3 NIV to a wall power supply.



The Airvo 3 NIV delivers reduced humidity in the breathing gas during Optiflow high flow therapy until the battery is nearly depleted, where humidity is turned off to maintain the delivery of flow and oxygen. During NIV therapy humidity is turned off to maintain the delivery of flow and oxygen.

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