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CPAP/Auto CPAP/BIPAP ST25 / ST30

User Guide



Product

The DS-5 is VentMed's Positive Airway Pressure (CPAP) device.

The DS-6 is VentMed's Auto-adjusting Pressure (Auto CPAP) device.

The DS-7 is VentMed's Bilevel Positive Airway Pressure ST25 (BIPAP ST25) device. The DS-8 is VentMed's Bilevel Positive Airway Pressure ST30 (BIPAP ST30) device.

CPAP mode provide same inspiratory and expiratory therapy pressure in one breathing cycle, Bilevel mode provide different inspiratory and expiratory therapy pressure.

WARNING

Read this entire guide before using the device.

In the US, Federal law restricts this device to sale by or on the order of a physician. Indications for use

The VentMed CPAP and Auto CPAP is indicated for the treatment of obstructive sleep apnea (OSA) in patients weighing more than 66 lb (30 kg). The humidifier is integrated in device.

The VentMed BIPAP ST25 and BIPAP ST30 are indicated for the treatment of obstructive sleep apnea (OSA)/ central sleep apnea (CSA)/ mixed sleep apnea (MSA) in patients weighing more than 66 lb (30 kg). The humidifier is integrated in device.

Adverse effects

You should report severe headache, unusual chest pain or increased breathlessness to your prescribing physician. An acute upper respiratory tract infection may require temporary discontinuation of treatment.

The following side effects may arise during the course of therapy with the device:

- $\hfill\square$ drying of the nose, mouth, or throat
- □ nosebleed
- □ bloating
- ear or sinus discomfort
- □ eye irritation

 \Box skin rashes.

Contraindications

Positive airway pressure therapy may be contraindicated in some patients with the following pre-existing conditions:

- pneumothorax
- □ dehydration
- □ severe bullous lung disease
- □ pathologically low blood pressure
- □ cerebrospinal fluid leak, trauma recent, or cranial surgery.

Packing list of the device

- Device with integrated humidifier
- Water tank
- Power supply unit
- Travel bag
- Measuring cup

Contact your care provider for a range of accessories available for use with the device including: Air tubing/Air filter/Mask

Use the mask/air tubing that complies with ISO17510/ISO80601-2-70/EN ISO10993.

ARNING: Failure to use a mask or accessory that minimizes rebreathing of carbon dioxide or permits spontaneous breathing can cause asphyxiation.

About device



- 1 Air outlet
- 2 SD card slot
- 3 Power inlet
- 4 Air filter cover
- 5 Water tank
- 6 Water tank open switch
- 7 Screen
- 8 Start/Stop button: Press to start/stop therapy

About the control panel

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Start/Stop button

Up button



Down button

Enter button

Press to start/stop therapy.

Press to select previous item.

Press to select next item.

Press to enter the selected item.

Different icons may be displayed on the screen at different times including:



Setup

User Guide



Do not overfill the water tank as water may enter the device and air tubing.

- 1. Place the device on a stable level surface.
- 2. Plug the power connector into the rear of the device. Connect one end of the power cord into the power supply unit and the other end into the power outlet.
- 3. Connect the air tubing firmly to the air outlet located on the rear of the device.
- 4. Open the water tank cover and fill it with distilled water up to the maximum water level mark. Do not fill hot water in the water tank.
- 5. Close the water tank cover.
- 6. Connect the free end of the air tubing firmly onto mask. For detailed information, read the mask user guide.
- 7. Physician should ensure the compatibility of the equipment and all of the parts and accessories used to connect to the patient before use; should ensure that the therapeutic pressure settings were determined for the patient individually with the configuration of the equipment to be used, including accessories; and should

periodically reassess the setting(s) of the therapy for effectiveness.

8. The equipment must not be covered or positioned in such a way that the operation or performance of the equipment is adversely affected.

Starting therapy

- 1. Fit your mask.
- 2. Press Start/Stop button.
- 3. You will know that therapy is on when the air fan icon is rota

The current treatment pressure is shown in white words.

During ramp time the pressure is gradually increasing and you will see a blue triangle icon on the screen once the prescribed pressure is reached.

The screen will go black automatically after a short period of time. You can press any button to turn it back on.

Stopping therapy

1. Remove mask.

2. Press Start/Stop button, therapy will stop automatically after a few seconds. The Sleep Report now gives you a summary of therapy session.



Compliance/Week Avg: H--Indicated the number of hours of therapy you received in one week.

H Today--Indicated the number of hours of therapy you received today. Compliance/Month: H--Indicated the number of hours of therapy you received in one month.

Sleep Report

1. In work interface, press Up or Down button to check sleep report date and waveform.



W M: Work Mode	Press: Therapy Pressure	T V: Tidal Volume
Insp T: Inspiratory Time	M V: Minute Volume	Exp T: Expiratory
Time		
Leak: Air Leakage	BPM: Breaths Per Minute	

My Options

The device has been set up for your needs by your care provider, but you may want to make small adjustments to make your therapy more comfortable. Highlight My Options and press Up/Down button to see current setting. You can personalize your options.

Ramp Time

Designed to make the beginning of therapy more comfortable, Ramp Time is the period during which the pressure increases from a low start pressure to the prescribed treatment pressure. You can set Ramp Time from 0 to 60minutes.

To adjust Ramp Time:



- 1. In Options, press enter button to highlight Ramp Time.
- 2. Press up/down button to adjust the ramp time to your preferred setting.
- 3. Press enter button to save the change.

Humidity Level

The humidifier moistens the air and is designed to make therapy more comfortable. Can set the Humidity Level to off (0) or between 1 and 5, where 1 is the lowest humidity setting and 5 is the highest humidity setting.

To adjust the Humidity Level:



- 1. In Options, press enter button to highlight Humidity Level.
- 2. Press up/down button to adjust humidity level.
- 3. Press enter button to save the change.

Work Mode

CPAP mode for DS-5 CPAP. CPAP, Auto mode for DS-6 Auto CPAP. CPAP, Auto, S, T, ST mode for DS-7 BIPAP ST25. CPAP, Auto, S, T, ST, APCV mode for DS-8 BIPAP ST30.

To Unlock Woke Mode parameters



- 1. Work Mode parameters will be locked automatically after setting in 15 minutes.
- 2. Long press enter button over 5 seconds to highlight the column to unlock. To adjust the working pressure

To adjust the working pressure



1. In work mode CPAP, press enter button to highlight Pressure. Press up/down button to adjust working pressure. You can set pressure 4-20 cmH2O for CPAP mode.

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 In work mode Auto CPAP, press enter button to highlight Pressure. Press up/down button to adjust Max and Min working pressure. You can set pressure 4-20 cmH2O for Auto CPAP mode.

Home>Options	Home>Options>Work Mode	> My Options > Work Mode
∢ Home	My Options) IPAP
Ramp Time 30 min	Work Mode S	10.0
Humidity Level 3	IPAP 10.0 cmH2O	10.0 cmH2O
Work Mode S >	EPAP 4.0 cmH2O	
Auto ON ON	Insp. Trigger 2	
Auto OFF ON	Exp. Level 2	5 10.0
Time Setting	Insp. Sens. 2	
		M S SUS M
Home>Options>Work Mode	Home>Options>Work Mode	> My Options > Work Mode
My Options) IPAP
Work Mode T	Work Mode T	10.0
IPAP 10.0 cmH2O	IPAP 10.0 cmH2O	10.0 cmH2O
EPAP 4.0 cmH2O	EPAP 4.0 cmH2O	
Backup Rate 10 bpm	Backup Rate 10 bpm	
I / E Rate 33 %	I / E Rate 33 %	> 10.0
Insp. Trigger 2	Insp. Trigger 2	
		M S SUS M
> My Options > Work Mode	Home>Options>Work Mode	> My Options > Work Mode
♦ My Options) IPAP
Work Mode ST	Work Mode ST	10.0
IPAP 10.0 cmHzO	IPAP 10.0 cmH2O	10.0 cmH2O
EPAP 40 cmHoG	EPAP 4.0 cmH2O	
Packup Data	Backup Rate 10 bpm	
	I / E Rate 33 %	10.0
Insp. /Exp. % 33%	Insp. Trigger 2	
Insp. Trigger 2		
A DESCRIPTION OF THE OWNER	Exp. Level 2	

3. In work mode S, T, ST, press enter button to highlight IPAP (Inspirate Positive Airway Pressure) . Press up/down button to adjust IPAP working pressure. You

can set pressure 4-25 cmH2O for DS-7 BIPAP ST25 and 4-30 cmH2O for DS-8 BIPAP ST30



 In work mode S, T, ST, press enter button to highlight EPAP (Expirate Positive Airway Pressure). Press up/down button to adjust EPAP working pressure. You can set pressure 4-20 cmH2O for DS-7 BIPAP ST25 and DS-8 BIPAP ST30.

Attention: Inspirate pressure must be higher than expirate pressure.

To adjust Backup Rate

Designed to set device inspirate and expirate therapy times in 1 minute. You can set 5-50bpm in mode T & ST.

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TO adjust I/E Rate

Designed to set device different inspirate and expirate therapy percentage in a whole therapy duration. You can set 10%-80% in mode T & ST.



TO adjust Inspirate Trigger

You can set 1-5 in mode S/T/ST, "1" is the most sensitive, "5" is the slowest.



TO adjust Expirate Level

You can set 1-5 in mode S & ST, "1" is the most sensitive, "5" is the slowest.

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TO adjust Inspirate Sensitivity

You can set 1-5 in mode S & ST, "1" is the most sensitive, "5" is the slowest.



To set APCV mode

In APCV mode, you can set IPAP/EPAP/Backup Rate/Inspirate Trigger/ Expirate Level/Inspirate Sensitivity, the adjust process is same as ST mode.

Home>Options	>Work Mode
My Options	
Work Mode	APCV
IPAP	25.0 cm+t2O
EPAP	10.0 cmH2O
Backup Rate	12 bpm
Insp. Duration	1.0 \$
Insp. Trigger	3
Exp. Level	3
Insp. Sens.	3
	- C M

To adjust Inspirate Duration



You can set from 0.5 seconds to 4 seconds.

TV. V. Switch



Designed to set target tidal volume ON or OFF.

User Guide

To adjust TV.V.



Designed to set target tidal volume, you can set from 200ml to 1200ml.

To adjust Max. IPAP



Designed to set maxium IPAP (Inspirate Positive Airway Pressure), you can set from 10 cmH2O to 30 cmH2O.

To adjust Min. IPAP



Designed to set minimum IPAP (Inspirate Positive Airway Pressure), you can set from 4 cmH2O to 20 cmH2O.

Auto ON



Designed to make the device start automatically in a while after you wear mask.

Auto OFF



Designed to stop the device work automatically in 10seconds after you take mask away.

Time Setting



Pressure Relief

The device already automatically set EPR (Expiratory Pressure Relief) function; you may find it easier to breathe out. This can help you get used to therapy. Can set the expiritory pressure relief Level to off (0) or between 1 and 3, where 1 is

the lowest expiratory pressure relief and 3 is the highest setting.



Caring for device

It is important that you regularly clean the device to make sure device work normally. The following sections will help you with disassembling, cleaning, checking and reassembling the device.

Disassembling

- 1. Open the water tank cover, take water tank away from the device.
- 2. Open the water tank and discard any remaining water.
- 3. Hold the cuff of the air tubing and gently pull it away from the device.

Cleaning

Clean the device weekly as described. Refer to the mask user guide for detailed instructions on cleaning your mask.

1. Wash the water tank and air tubing in warm water using mild detergent. Do not wash in a dishwasher or washing machine.

- 2. Rinse the water tank and air tubing thoroughly and allow to dry out of direct sunlight and/or heat.
- 3. Wipe the exterior of the device with a dry cloth.

Checking

Regularly check the water tank, air tubing and the air filter for any damage.

- 1. Check the water tank:
- □ Replace it if it is leaking or has become cracked, pitted or cloudy.
- \Box Replace it if the seal is cracked or torn.
- Remove any white powder deposits using a solution of one part household vinegar to 10 parts water.
- 2. Check the air filter and replace it at least every six months. Replace it more often if there are any holes or blockages by dirt or dust.

To replace the air filter:



- □ Open the air filter cover and remove the old air filter. The air filter is not washable or reusable.
- □ Place a new air filter then close the air filter cover.
- □ Make sure the air filter is fitted at all times to prevent water and dust from entering the device.
- 3. Check the air tubing and replace it if there are any holes, tears or cracks.

Reassembling

When the water tank and air tubing are dry, can begin to reassemble the parts.

- 1. Connect the air tubing firmly to the air outlet located on the rear of the device.
- 2. Fill the water tank with distilled room temperature water up to the maximum water level mark.
- 3. Put water tank into the device. Close the water tank cover.
- 4. Connect the free end of the air tubing firmly onto the mask.

Therapy data

The device records therapy data for you and your care provider so they can view and make changes to your therapy if required. The data is recorded via a SD card. You can send SD card to your care provider by mail. When instructed by your care provider, remove the SD card.

Do not remove the SD card from the device when the device is working.

To remove the SD card:

- 1. Stop the device.
- 2. Push in the SD card to release it. Remove the SD card from the device.

Place the SD card in the protective folder and send it back to your care provider. Note: The SD card should insert in correct, should not be used for any other purpose.



Traveling

You can take this device with you wherever you go. Just keep the following in mind:

- □ Empty the water tank and put it into the device.
- Make sure you have the appropriate power cord for the region you are traveling to. For information on purchasing, contact your care provider.
- □ Use the travel bag provided to prevent damage to the device.
- If you are using an external battery, should turn off the humidifier in order to maximize the life of your battery. Do this by turning the Humidity Level to Off.

Do not use the device with water in the water tank on a plane due to the risk of inhalation of water during turbulence.

Troubleshooting

If you have any problems, have a look at the following troubleshooting topics. If you are not able to fix the problem, contact your care provider or VentMed. Do not try to open the device.

Problem/possible cause	Solution
Air is leaking from around mask Mask may be fitted incorrectly.	Make sure mask is fitted correctly and firmly. Read your mask user guide for detailed information.
Air pressure in mask seems too high (it feels like getting too much air) Ramp may be turned off.	Use the Ramp Time option.
Air pressure in mask seems too low (it feels like I am not getting enough	
air) Ramp may be in progress.	Wait for air pressure to build up or turn Ramp Time off.
My water tank is leaking	
Water tank may not be assembled	Check for damage and reassemble the
correctly.	water tank correctly.
Water tank may be damaged or cracked.	Contact your care provider for a replacement.
Nose is feeling dry or blocked	
Humidity level may be set too low.	Adjust the Humidity Level.
Screen is black	
Backlight on the screen may have	Press any button to turn it back on.
turned off. It turns off automatically	Connect the power supply and make sure
after a short time.	the plug is fully inserted.
Power may not be connected.	
Have stopped therapy, but the device	Device blows a small amount of air in
is still blowing air	order to avoid condensation in the air
Device is cooling down.	tubing. It will stop automatically after 20
	minutes.

Problem/possible cause	Solution
High leak detected, check water tank, tank seal or water tank cover	
Water tank may not be inserted properly.	Make sure the water tank is correctly inserted.
Water tank seal may not be inserted properly.	Open the water tank and make sure that the seal is correctly inserted.
Droplets of water on nose, in the mask and air tubing	
Humidity level may be set too high.	Adjust the Humidity Level.
Mouth is very dry and	Increase the Humidity Level.
uncomfortable	You may need a chin strap to keep your
Air may be escaping through mouth	mouth closed or a full face mask.
High leak detected, connect your	Make sure the air tubing is firmly
tubing	connected at both ends.
Air tubing may not be connected properly.	Make sure mask is fitted correctly. See your mask user
Mask may be fitted incorrectly.	guide for fitting instructions or use the Mask Fit function to check your mask fit and seal.
Tubing blocked, check your tubing	Check the air tubing and remove any
Air tubing may be blocked.	blockages.
SD card error	
SD card may not be inserted correct	ly. Power off the device, remove and reinsert the SD card.
Air filter blocked	Check the air filter and replace it if there are any blockages.
Water in the air tubing	Empty the water from the air tubing. Disconnect the power
	supply and then reconnect it to restart the device.
All other error messages	Contact your care provider. Do not open the device.

Reassembling parts

Water tank is designed to easily come off in order to avoid damage to the parts or the device. You can easily reassemble them as described below.



To insert the water tank seal:

- 1. Place the seal into bottom water tank.
- 2. Press down along all edges of the seal until it is firmly in place.

To reassemble the water tank:

1. Insert both up water tank into bottom water tank and sealed both sides.

General warnings and cautions

MWARNING

- □ Make sure that you arrange the air tubing correctly so that it will not twist around the head or neck.
- □ Keep the power cord away from water source or hot surfaces.
- Make sure the power cord and plug are in good condition and the device is not damaged.
- Do not open or modify the device. Repairs and servicing should only be performed by your care provider.
- Beware of electrocution. Do not immerse the device, power supply or power cord in water. If liquids are spilled into or onto the device, unplug the device and let the parts dry. Always unplug the device before cleaning and make sure that all parts are dry before plugging it back in.
- □ If you notice any unexplained changes in the performance of the device, if it is making unusual sounds, if the device or the power supply are dropped or

mishandled, or if the enclosure is broken, discontinue use and contact your care provider.

- Supplemental oxygen must not be used while smoking or in the presence of an open flame.
- Always make sure that the device is turned on and airflow generated before the oxygen supply is turned on. Always turn the oxygen supply off before the device is turned off, so that unused oxygen does not accumulate within the device enclosure and create a risk of fire.
- Sources of oxygen must be located more than 1 m from the equipment to avoid the risk of fire and burns.
- Do not perform any maintenance tasks while the device is in operation.

- Use only compliant parts and accessories with the device. Non-compliant parts may reduce the effectiveness of the treatment and/or damage the device.
- Be careful not to place the device where it can be bumped or where someone is likely to trip over the power cord.
- Use only VentMed masks recommended by VentMed or by the prescribing doctor with this device. Fitting the mask without the device blowing air can result in rebreathing of exhaled air. Make sure that the mask vent holes are kept clear and unblocked to maintain the flow of the fresh air into the mask.
- □ Keep the device dry, clean and clear of anything (eg, clothes or bedding) that could block the air inlet or cover the power supply unit.
- Do not place the device on its side as water might get into the device.
- Do not block the air tubing and/or air inlet of the device while in operation as they could lead to overheating of the device.
- Incorrect system setup may result in incorrect mask pressure reading. Ensure the system is correctly set up.
- □ Make sure that the water tank is empty before transporting the device.
- □ When use the humidifier, please always place the device on a level surface lower than your head to prevent the mask and air tubing from filling with water.
- □ Leave the water tank to cool for ten minutes before handling to allow the water to cool and to make sure that the water tank is not too hot to touch.
- Do not use bleach, chlorine, alcohol, or aromatic-based solutions, moisturizing or antibacterial soaps or scented oils to clean the device, the water tank or air tubing. These solutions may cause damage or affect the humidifier

performance and reduce the life of the products.

 \Box Periodically reassess the setting(s) of the therapy for effectiveness.

Technical specifications

Туре	Item	Specifications
60W power supply	AC input range	100–240V~, 50–60Hz,
unit (Only use		2A maximum, Class II
with IEC60601-1	DC output	24V2.5A
approved adaptor)		
	Operating temperature	+41°F to +95°F (+5°C to +35°C)
		Note: The air flow for breathing produced by this therapy
		device can be higher than the
Environmental		temperature of the room.
conditions		Under extreme ambient
		temperature conditions
		(104°F/40°C) the device
		remains safe.
5	Operating humidity:	10 to 95% relative humidity,
8		non-condensing
	Operating altitude	Sea level to 8,500' (2,591 m);
		air pressure range 1013 hPa to
		738 hPa
	Storage and transport temperature	-4°F to +140°F (-20°C to +60°C)
	Storage and transport humidity	y 5 to 95% relative humidity,
e		non-condensing
Electromagnetic	The VentMed device complies	
compatibility	with all applicable	
	electromagnetic compatibility	
	requirements (EMC) according	9
	to IEC60601-1-2:2014, for	
	residential, commercial and	

		Ventitied
	light industry environments	
	Class II (double insulation),	
60601-1:2005+A1 2012 classification	Type BF, Ingress protection IPX1.	
Sensors	Pressure sensor	Internally located at device outlet, analog gauge pressure type, 0 to 40 cm H2O
	Flow sensor	Internally located at device outlet, analog gauge flow type, 50 to +200 L/min
Maximum single	Device will shut down in	
fault steady	the presence of a single fault i	f
pressure	the steady state pressure	
	exceeds: 35 cm H2O for more	
	than 1 sec.	
Sound	Pressure level measured	53 dBA with uncertainty of
	according to ISO 80601-2-70	2 dBA under background
	dual number noiseemission	A weighted sound power leve
	values in accordance with IS	O is 61dba under backgrou
	4871-1996	sound level 36dBA
	Dimensions (H x W x D) [.]	280 mm x 140 mm x 95 mm
1	Air outlet (complies with ISO	22 mm
	5356-1:2004)	
Physical-device	Weight (device and water tank	() 1.6kgs
and water tank	Water capacity	To maximum fill line
		220 ML, minimum 100ML
	Water tank - material	Injection molded plastic
		and silicone seal
	Maximum heater plate	68°C
Temperature	Cut-out	102°C

		VCITUNCU	
	Maximum gas temperature	41°C	
Operating pressure DS-5 CPAP		4 to 20 cm H2O	
range(IPAP)	DS-6 Auto CPAP	4 to 20 cm H2O	
	DS-7 BIPAP ST25	4 to 25 cm H2O	
	DS-8 BIPAP ST30	4 to 30 cm H2O	
Operating pressur	e DS-5 CPAP		
range(EPAP)	DS-6 Auto CPAP	4 to 20 cmH2O	
	DS-7 BIPAP ST25		
0	DS-8 BIPAP ST30		
Air Filter		Material: Polyester non	
		woven fiber	
		Average arrestance: >75%	
		for ~7 micron dust	
	Device, power supply unit	5 years	
Design life	Water tank, air tubing	6 months	
Nataa			

Notes:

Manufacturer will provide circuit diagrams, component part lists, descriptions, calibration instructions to assist to service personnel in parts repair.

- Manufacturer reserves the right to change specifications without notice.
- The temperature and relative humidity settings displayed are not measured value Do not use electrically conductive or antistatic air tubing.



User Guide

Displayed values

Value	Range			Display resolution	
Pressure sensor at a	e sensor at air DS-5 CPAP			0.1 cmH2O	
outlet: Mask pressure	DS-6 Auto CPAP			0.1cmH2O	
	DS-7	BIPAP	ST25:	4–25	0.1 cmH2O
	cmH2()			0.1 cmH2O
	DS-8	BIPAP	ST30:	4-30)
	cmH2O				

Value Accuracy

/			
Value	Accuracy		
Static Airway Pressur	e ±0.5cmH2O + 4% of the set pressure		
measurement: Mask			
pressure			
Dynamic Airway	±3cmH2O + 5% of the set pressure		
Pressure			
measurement(Both	The percentage of each inspiratory and expiratory p	nase	
Cpap/BIPAP): Mask	to be taken into the calculation for determining t	he	
pressure	accuracy is (I/E=1:1)		
Remark: Results are expressed at ATPD (Ambient Temperature and Pressure			
Dry).			

Environmental information



This device should be disposed of separately, not as unsorted municipal waste. To dispose of your device, you should use appropriate collection, reuse and recycling systems available in your region. The use of these collection, reuse and recycling systems is designed to reduce pressure on natural resources and prevent hazardous substances from damaging the environment.

If you need information on these disposal systems, please contact your local waste administration. The crossed-bin symbol invites you to use these disposal systems. If you require information on collection and disposal of the device please contact your care provider or VentMed.

Symbols

The following symbols may appear on the product or packaging.

	Indicates a warning or caution
SN	Serial number
*	Type BF applied part
	Maximum water level
$\land \land$	Caution! Hot Surface. Do not touch.
IPX1	Water proof Grade 1
	Manufacturer
EC REP	European Authorized Representative
	Environmental information
ī	Follow instructions before use
	Direct current
+5°C +35°C	Temperature limitation
<u>%</u>	Humidity limitation
\sim	Manufacturer Date

Limited warranty

VentMed Ltd warrants that your VentMed product shall be free from defects in material and workmanship from the purchase date for the period specified below.

Product	Warranty period
Humidifier water tanks	90days
CPAP, BIPAP devices (including	g 2 years
external	
power supply units)	
Humidifier	2 years

This warranty is only available to the initial consumer. It is not transferable.

If the product fails under conditions of normal use, VentMed will repair or replace, at its option, the defective product or any of its components.

This Limited Warranty does not cover:

a) Any damage caused as a result of improper use, abuse, modification or alteration of the product;

b) Repairs carried out by any service organization that has not been expressly authorized by VentMed to perform such repairs;

c) Any damage or contamination due to cigarette, pipe, cigar or other smoke.

Warranty is void on product sold, or resold, outside the region of original purchase. Warranty claims on defective product must be made by the initial consumer at the point of purchase.

This warranty replaces all other expressed or implied warranties, including any implied warranty of merchantability or fitness for a particular purpose. Some regions or states do not allow limitations on how long an implied warranty lasts, so the above limitation may not apply to you.

VentMed shall not be responsible for any incidental or consequential damages claimed to have resulted from the sale, installation or use of any VentMed product.

Further information

If you have any questions or require additional information on how to use the device, contact your care provider.

Technical

Description

Concerning

Electromagnetic

Immunity

Table 3: Guidance & Declaration - electromagnetic immunity

Guidance & D	Guidance & Declaration — electromagnetic immunity			
The models [DS-5, DS-6, DS	S-7, DS-8 are inten	ded for use in the electromagneti	
environment s	specified below.	The customer or t	he user of the models DS-5, DS-	
DS-7, DS-8 sl	nould assure tha	t It is used in such a	an environment.	
Immunity	IEC 60601 test	Compliance level	Electromagnetic environment	
test	level		- guidance	
Electrostatic	±8 kV contact	±8 kV contact	Floors should be wood,	
discharge	±2 kV, ±4 kV,	±2 kV, ±4 kV,	concrete or ceramic tile. If	
(ESD)	±8 kV,	±8 kV, ±15 kV air	floors are covered with	
IEC	±15 kV air		synthetic material, the relative	
61000-4-2			humidity should be at least 30	
Electrical fast	±2kV for	±2kV for power	Mains power quality should be	
transient/burs	power supply	supply lines	that of a typical commercial or	
t	lines	±1kV for	hospital environment.	
IEC	±1 kV for	interconnecting		
61000-4-4	Input/output	cable		
2	lines			
Surge	±1 kV line to	±1 kV line to	Mains power quality should be	
IEC	line	line	that of a typical commercial or	
61000-4-5	±2 kV line to		hospital environment.	
	earth			
Voltage dips,	<5 % UT	<5 % UT	Mains power quality should be	
short	(>95% dip in	(>95% dip in UT.)	that of a typical commercial or	
interruptions	UT.)	for 0.5 cycle	hospital environment. If the	
and	for 0.5 cycle	40 % UT	user of the models DS-5, DS-6,	
voltage	40 % UT	(60% dip in UT)	DS-7, DS-8 require continued	
variations on	(60% dip in	for 5 cycles	operation during power mains	
power supply	UT)	70% UT (30% dip	interruptions, it is	

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input	for 5 cycles	in UT)	recommended that the models
lines	70% UT	for 25 cycles	DS-5, DS-6, DS-7, DS-8 be
IEC	(30% dip in	<5% UT (>95 %	powered from an
61000-4-11.	UT)	dip in	uninterruptible power supply or
	for 25 cycles <5% UT (>95 % dip in UT) for 5 sec	UT) for 5 sec	a battery.
Power frequency (50- 60 Hz) magnetic field IEC 61000-4-8	30 A/m	30 A/m	Power frequency magnetic fields should be at levels characteristic of a typica location in a typical commercial or hospital environment.
NOTE UT is t	he a.c. mains vo	Itage prior to application	ation of the test level.

Annex A: Electromagnetic Compatibility Information

Guidance and manufacturer's declaration - electromagnetic emissions- this device is intended for use in the electromagnetic environment specified below. The

user of the device should ensure that it is used in such an environment.

Emissions Test	Complianc	Electromagnetic environment-guidance
	е	
		The device users RF energy only for its
RF emissions		internal function. Therefore its RF emissions
CISPR11	Group 1	are very low and are not likely to cause any

		interference in nearby electronic equipment.	
RF emissions	Class B	This device is suitable for use in all	
CISPR11		establishments, including domestic	
Harmonic emission	s Class A	establishments and those directly connected	d to
IEC61000-3-2		the public low-voltage power supply network	K
Voltage fluctuation	Conformity	/ that supplies buildings used for domestic	
/flicker emissions		purposes.	
IEC61000-3-3			
			i

Guidance and manufacturer's declaration - electromagnetic immunity – this device is intended for use in the electromagnetic environment specified below. The user of the device should make sure that it is used in such an environment.

Immunity test	IEC60601 test	Compliance level	Electromagnetic
	level		environment-guidance
			Floor should be wood,
Electrostatic	±6KV contact	±6KV contact	concrete or ceramic tile. If
discharge	±8KV air	±8KV air	floors are covered with
(ESD)			synthetic material, the
IEC61000-4-			relative humidity should be
2			at least 30%.
Electrical fast	±2KV for	±2KV for	Mains power quality
transient /	power supply lines	power supply lines	should be that of a typical
burst	±1KV for	±1KV for	home or hospital.
IEC61000-4-	input/output lines	input/output lines	

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	±1K	±1KV	Mains power quality
Surge	differential mode	differential mode	should be that of a typical
IEC61000-4-	±2KV	±2KV	home or hospital.
5	common mode	common mode	

Immunity test	IEC60601 test level	Compliance level	Electromagnetic
			environment-guidance
	<5% U⊤	<5% U⊤	Main power quality
Voltage dips,	(>95% dip in U $_{ op}$),	(>95% dip in U $_{ au}$),	should be that of a
short	for 0.5 cycle	for 0.5 cycle	typical commercial or
interruptions			hospital environment. If
and voltage	40% U⊤(60% dip in	40% U⊤(60% dip in	the use of the device
variations on	U⊤), for 5 cycles	U⊤), for 5 cycles	requires continued
power supply			operation during power
input lines	70% U⊤	70% U⊤	mains interruptions, it is
IEC61000-4-11	(30% dip in U⊤)	(30% dip in U⊤)	recommended that the
	for 25 cycles	for 25 cycles	device be powered form
			an uninterruptible
	<5% U⊤	<5% U⊤	power supply or from a
	(>95% dip in U⊤)	(>95% dip in U⊤)	battery.
	for 5 s	for 5 s	
Power			Power frequency

User Guide			VentMed
frequency	3A/m	3A/m	magnetic field shall be
(50/60Hz)			typical level of power
magnetic field			frequency magnetic
IEC61000-4-8			fields in hospital or
			home environment.

Note: U_T is the AC mains voltage prior to application of the test level.

Guidance and manufacturer's declaration - electromagnetic immunity - this

device is intended for use in the electromagnetic environment specified below. The

user of the device should make sure that it is used in such an environment.

Immunity	IEC60601 test	Compliance	e Electromagnetic
test	level	level	environment-guidance

User Guide			VentMed	
(-778)			Portable and mobile RF	
			communications equipment should	
			be used no closer to any part of the	
			device, including cables, than the	
			recommended separation distance	
			calculated from the equation	
			application to the frequency of the	
			transmitter.	
			Recommended separation distance	
			d=1.2 √p	
Conducted			d=1.2√p 80MHz to 800MHz	
RF	3Vrms	3Vrms	d=2.3√p 800MHz to 2.5GHz	
IEC61000-4	150kHz to			
-6	80MHz			

Immunity test	IEC60601 test	Compliance	Electromagnetic
	level	level	environment-guidance
			Where P is the
Radiated RF	3V/m	3V/m	maximum output
IEC61000-4-3	80MHz to		power rating of the

User Guide		 VentMed
	2.5GHz	transmitter in watts
		(W) according to the
		transmitter
		manufacturer and d is
		the recommended
		separation distance in
		meters (m).
		Field strengths from
		fixed RF transmitter,
		as determined by an
		electromagnetic site
		survey, should be less
		than the compliance
		level in each
		frequency range.

Immunity test	IEC60601 test	Compliance	Electromagnetic
	level	level	environment-guidance
			Interference may
			occur in the vicinity of
			equipment marked

with the following symbol:

Note 1: At 80MHz, the higher frequency range applied.

Note 2: These guidelines may not be applied in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

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

b. Over the frequency range 150kHz to 80MHz, the field strengths should be less than 3 V/m.

Maximum flow rate

1. CPAP

Pressure(cmH2O) Maximum flow rate(L/Min)
4	77
8	111
12	140

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16	158
20	164

2. BIPAP

Pressure(cmH2O)	Maximum flow rate(L/Min)
4	77
9	119
15	156
20	164
25	164
30	164

Manufacturer:

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