

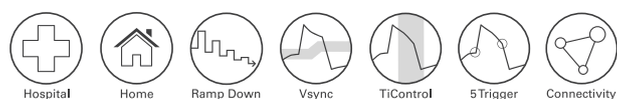
Lumis™ 100

VPAP S

Personalised ventilation made easy

Lumis 100 VPAP S is a noninvasive ventilator designed for spontaneously breathing non-dependent patients with respiratory insufficiency.

It's easy to set up and use, and features built-in wireless connectivity to AirView™ – ResMed's cloud-based patient management system – for remote monitoring.





Individually responsive with IntelligentAir

IntelligentAir is a collection of ResMed technologies that can tailor therapy to individual breathing needs. Automatically adapting to changes during the night and at different stages of a patient's condition, IntelligentAir makes personalised ventilation possible.



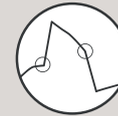
Vsync

Vsync provides excellent patient-ventilator synchrony, even in the presence of significant leak.



TiControl

TiControl™ lets you set min and max limits on either side of the patient's ideal inspiratory time to encourage spontaneous breathing.



5Trigger

Trigger and cycle helps you optimise settings according to the patient's condition, using five trigger and cycle sensitivity levels.

Key features

QuickNav for low-touch therapy adjustment

By simply double-clicking the home button on the device, you can quickly and easily toggle between the Settings and Monitoring screens.

ClimateControl Auto for automatic humidification

When used with a HumidAir™ heated humidifier and ClimateLineAir™ heated tube, Lumis delivers humidification automatically – no settings to change and no complicated menus to navigate. So you can set your patients up to receive all the benefits of humidification as soon as they turn on their device.

Ramp and Ramp Down for extra comfort

Lumis offers both Ramp and Ramp Down features to help patients ease into and out of each therapy session. Ramp reaches the prescribed therapy level gradually and comfortably, while Ramp Down gradually reduces the pressure support and EPAP to help ease patients off therapy.

AirView for remote patient management

Your patients' therapy data can be sent seamlessly and securely to AirView, where you can view it remotely, and gain timely access to the most recent therapy statistics and trend data. AirView's 'remote assist' and 'remote settings' changes also allow for easy and efficient device management.

Product codes

Lumis 100 VPAP S with HumidAir heated humidifier, ClimateLineAir heated tube and built-in wireless connectivity **28008**

Note: Technical specifications may change without notice. Always refer to your device user guide.

Technical specifications

Modes	CPAP, S
Operating pressure range	2–25 cm H ₂ O (2–25 hPa) in S mode 4–20 cm H ₂ O (4–20 hPa) in CPAP mode
Sound pressure level	26.6 dBA with uncertainty of 2 dBA as measured according to ISO 17510-1:2009 (CPAP mode)
Dimensions (H x W x D)	116 mm x 205 mm x 150 mm (Device only) 116 mm x 255 mm x 150 mm (Device with HumidAir humidifier)
Weight	1106 g (Device only) 1268 g (Device with HumidAir humidifier)
90W power supply unit	Input range: 100–240V, 50–60Hz, 57VA (typical power consumption) 108VA (peak power consumption)
Operating temperature humidity/altitude	+5°C to +35°C, 10–95% relative humidity, non-condensing/sea level to 2,591 m; air pressure range 1013 hPa to 738 hPa
Storage and transport temperature/humidity	-20°C to +60°C/5–95% relative humidity, non-condensing
Housing construction	Flame retardant engineering thermoplastic
Supplemental oxygen	Recommended maximum flow: 15 L/min (CPAP, S)
Standard air filter	Polyester non-woven fibre
Tubing	SlimLine™ and ClimateLineAir (15 mm); Standard (19 mm)
Air outlet	22 mm air outlet complies with ISO 5356-1:2004
Electromagnetic compatibility	Requirements (EMC) according to IEC60601-1-2: 2007 for residential, commercial, and light industry environments
Aircraft use	ResMed confirms that device meets the Federal Aviation Administration (FAA) requirements (RTCA/DO-160, section 21, category M) for all phases of air travel
IEC 60601-1: 2006 classification	Class II (double insulation), Type BF Ingress protection IP22



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