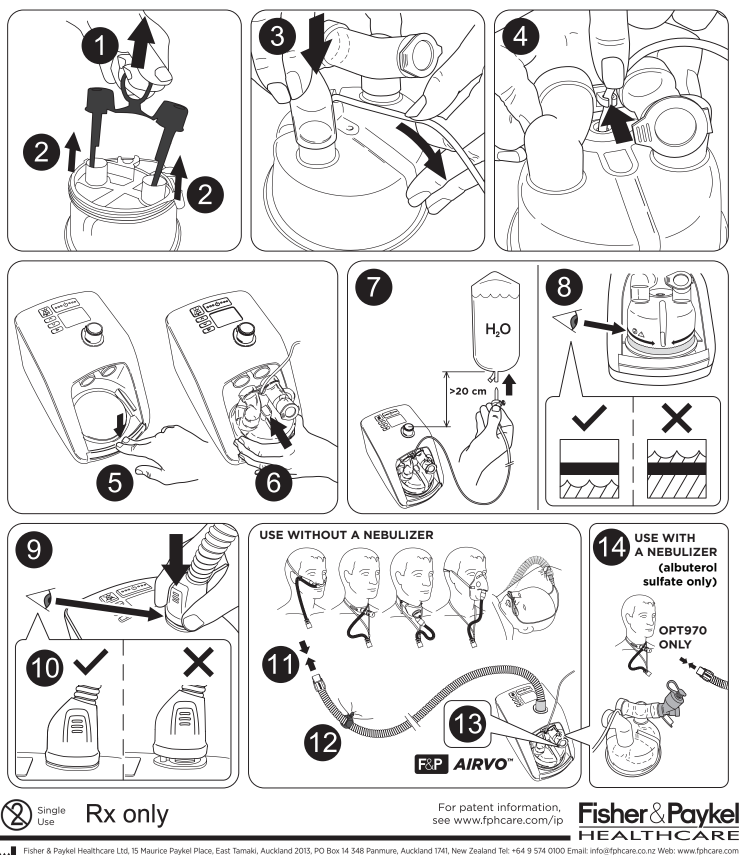


Tube and Chamber Kit

(with nebulizer adapter)

900PT563



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Airvo™ Tube and Chamber Kit with nebulizer adapter - 900PT563	Performance:			
Indications for use - Airvo nebulizer adapter: When used with a nebulizer:	Albuterol sulphate (1 mg/mL, 2.5 mL)	10 L/min	20 L/min	30 L/min
When used with a nebulizer: The nebulizer adapter is a medical device accessory for single-patient use to facilitate aerosolization of Albuterol sulfate for inhalation to adult patients receiving high-flow	Total delivered dose (µg)*	1362.9 - 2087.7	548.6 - 1938	383.3 - 1461.8
humidified breathing gases via tracheostomy patient interface. Intended for use by healthcare professionals in hospitals or long-term care facilities.	Total respirable dose (µg, 1-5µm)*	1035.9 - 1550.3	470.7 - 1428.2	387.9 - 837.3
When used without a nebulizer: For use in hospitals and long-term care facilities, for the delivery of humidified	Respirable fraction (%)	63 - 87.1	59.3 - 93.8	31.3 - 100
respiratory gases to patients via nasal, tracheostomy and mask interfaces.	Coarse particle dose (µg, >4.7µm)*	206.6 - 528.2	0 - 507.8	0 - 531.9
System Specifications: Circuit Length: 1.8 m (6 ft) + interface.	Coarse particle fraction (>4.7µm)	14.9 - 27.6	4.7 - 32.4	0 - 47.2
Use WITHOUT a nebulizer	Fine particle dose (µg, <4.7µm)*	1111.3 - 1604.5	519 - 1500.8	448.5 - 987.4
 If never used with a nebulizer, the 900PT563 kit must be disposed of after a maximum of 14 days' use. 	Fine particle fraction (<4.7µm)	72.4 - 85.1	67.6 - 95.4	52.8 - 100
Do not soak, wash or sterilize. Setup:	Mass-Median Aerosol Diameter (MMAD) (μm)*	2.67 - 2.94	2.41 - 3.05	2.08 - 3.00
 Airvo 2 humidifier ("Airvo") on a flat surface. For use at flows from 2 to 60 L/min depending on the patient interface. OPT316/OPT318/OPT942/OPT946/OPT1042/OPT1044/OPT1046 nasal interfaces OR OPT970 tracheostomy interface OR OPT980 mask adapter interface 	Geometric Standard Deviation (GSD) (µm)*	1.56 - 2.30	1.18 - 2.58	0.1 - 4.91
<u>WARNING</u> : Use of a non-approved accessory could impair performance or compromise safety.	* 95% confidence intervals			
Steps 1 - 7: Assemble and connect water chamber.	Testing carried out with one Aerogen Solo nebulizer, three sets of Airvo 2, 900PT563 kit and OPT970 interface and three tests per set. Cascade impactor testing conducted at 15 L/ min using NGI.			
Step 8: Ensure water level is rising but not going over the maximum water level.				
Incorrect water level Replace MR290 chamber	 WARNINGS: For single patient use only. Reuse may r Attempting to reprocess will result in de 	result in transmis gradation of ma	sion of infectious terials and rende	substances. r the product
Use USP sterile/distilled water for inhalation, or equivalent.	 defective. Using the breathing tube for longer than the specified time can result in serious injury, including infection. Do not use the auto-fill MR290 chamber if it has been dropped or been allowed to run dry this could lead to the chamber over-filling. Do not use the MR290 chamber if the water level rises above the maximum water level line this may lead to water entering the patient's airway. Never operate the unit if the breathing tube has been damaged with holes, tears or kinks. Do not block the flow of air through the unit and breathing tube. 			
Step 9: Connect breathing tube to Airvo.				
Step 10:				
Make sure blue connector is fully located into place. Step 11:				
Connect breathing tube to patient interface once the system has warmed up. Position breathing tube below patient interface so that condensate flows away from the patient. Step 12:	 California residents please be advised o product contains chemicals known to th and other reproductive harm. For more 	f the following, p ne State of Califo	ursuant to Propo rnia to cause can	cer, birth defects
Connect breathing tube clip to patient clothing or bedding. Note: connecting to patient clothing may not be suitable for all patient groups.	 To avoid burns: Do not allow the breathing tube to remain of time. 	ain in direct cont	act with skin for p	orolonged periods
Step 13: In the absence of a nebulizer, make sure the nebulizer port plug is firmly in place.	 Do not use in the presence of a naked flame, to avoid fires. Do not add heat to any part of the breathing tube e.g. covering with a blanket, or heating it 			
<u>WARNING:</u> Loss of therapy may occur if the nebulizer port plug is not in place.	an incubator or overhead heater for a n			
Use WITH a nebulizer If used with a nebulizer at any time, the 900PT563 kit must be disposed of after a maximum of 7 days' use, whether nebulization is continuous or intermittent during this time. Do not soak, wash or sterilize. CAUTION: Using the kit with a nebulizer for longer than 7 days may cause damage to the	Cautions: • Avoid product contact with chemicals, • Use only USP sterile/distilled water for i • Nebulized medications may be emitted Condensate containing nebulized medi personal protective equipment such as	nhalation, or equ into the surroun cation may be pr	ivalent. ding area.	
kit, particularly the water chamber, and eventually the Airvo itself. Setup:	 To prevent condensation: Use in a room warmer than 18 °C (64 °F 			
WARNING: Nasal delivery of nebulized drugs for lung deposition is not approved by FDA. WARNING: Use of a non-approved accessory could impair performance or compromise	Remove/minimize the impact of anythir air conditioning, open window. To manage excessive condensation:	ng that may cool	the heated breat	hing tube, e.g. a fa
safety. Airvo 2 humidifier ("Airvo") on a flat surface. For use at flows from 10 to 30 L/min using the OPT970 patient interface. OPT970 tracheostomy interface Aerogen Solo nebulizer	 Place the Airvo below patient head heic Drain condensate back into the water cl necessary to first reduce the target flow drains safely and effectively. 	hamber. At highe v setting to 30 L/	min, to ensure th	
Compatible medications:	 Disconnect the patient interface from the Lift the patient end of the heated breath water chamber 	hing tube, allowir	ng tube. ng the condensat	e to run into the
This kit is compatible with the following medications only: Albuterol sulfate	water chamber.			
 Refer to drug manufacturers' instructions for correct usage and restrictions. <u>WARNING</u>: Do not nebulize the following medications, as they are incompatible with Airvo and its consumables and may cause patient harm: Alcohol-based medications 				
 Significantly acidic or basic medications eg. epoprostenol (Flolan) Saline >0.9% 				
 Step 14 (Nebulizer use): Follow Steps 1-13. Remove the nebulizer port plug from the nebulizer port and connect the nebulizer in 				
its place. <u>CAUTIONS:</u> • Refer to the nebulizer manufacturer's instructions for setup and usage information. • Do not add oxygen or other gases through the nebulizer port or use a flow-				
generating nebulizer (eg. jet nebulizer) as this will cause discrepancies in humidity, flow and oxygen outputs. • Ensure the Airvo is switched on with humidified gas flowing before the nebulizer is used.				
 The medication delivery efficiency decreases as the Airvo flow increases. Ensure that the patient receives adequate flow at all times when adjusting flow during nebulization. Titrate medication delivery as required. Nebulized medications can form residue and cause cosmetic discoloration of the breathing tube and chamber during normal operation. 				
 Nebulization may cause increased levels of condensate in the breathing tube and chamber. 				
Avoid skin contact with condensate as it may contain nebulized medication.				