

BREATHING SYSTEMS TECHNICAL DATA SHEET

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Release Date	09.08.2023		
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BREATHING CIRCUIT NAME/ REFERENCE NUMBER	Anesthesia Circuit, Adult, Extendible Tubing, I	atex-free Bag/31305020-15			
MANUFACTURER NAME	R VENT Medikal Uretim A.S. Tel: +90 232 853 9500				
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	35860- Izmir, Turkey				
REGULATORY APPROVALS AND	ISO 13485 – 31816401				
CERTIFICATION	CE Certificate – 2195-MED-1816401				
CLASSIFICATION	Disposable Medical Device				
	MDD 93/42/EEC				
	Class IIa Rule 2 Annex V, Article 3				
	Airiex V, Airicle 3				
GMDN CODE/DESCRIPTION	37704				
	Anaesthesia breathing circuit, single-use				
		medical gases from an anaesthesia unit/workstation to a patient			
		ed) during general anaesthesia. It includes breathing tubes and a Y-			
		entilation bag and appropriate connectors, and may include a carbon valve, or adjustable pressure limiting (APL) valve. This is a single-use			
	device.	valve, or adjustable pressure limiting (APL) valve. This is a single-use			
EMDN CODE/DESCRIPTION	R02010101				
EMBR CODE, BESCHII HOR	Breathing Circuits, w/o Water Trap				
FEATURES	Disposable breathing circuits may help reduce cross-contamination.				
	Available in a wide variety of tubing styles, components and configurations to meet specific needs.				
INTENDED USE	Disposable breathing circuit for conduction of respiratory gases between anesthesia machine or ventilator and				
	patient and intended for single use only. Sterile and Non-sterile options are available.				
	Breathing bag with connection hose (limb) intended for use with anesthesia delivery systems as a reservoir during				
	automatic ventilation and as a manual breath	ing bag during manual ventilation.			
TECHNICAL SPECIFICATIONS	Drawing:				
	22mmF 2 180cm 3 4				
	7 .				
	' <u> </u>				
	22mmF				
	7 180cm				
	2,0 liter				
	Latex-Free Bag				
	22mmM 22mmF				
	8	ů ,			
	Materials:				
	Components	Materials			
	1 22M – 22F Straight Connector	Ethylene vinyl acetate (EVA)			
	2 22 Mm Extendible Tubing 180cm	Polyvinyl Chloride (PVC) (PHT FREE)			
	3 Y Connector W/Out Port	Polypropylene (PP)			
	4 Tethered Cap	Low-density polyethylene (LDPE)			
	5 Elbow Connector with CO2 Port	Polypropylene (PP)			
	6 22MM Extendible Tubing	Polypropylene (PP)			
	7 22M-22M/15F Straight Connector	Polypropylene (PP)			
	8 2lt Breathing Bag	Neoprene			
	9 Long Connector Red Cap	Ethylene vinyl acetate (EVA)			
	This product does not contain any metallic parts.				





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	Anneanance As shows an discussion				
	Appearance: As shown on drawing				
	Recommended Patient: Adult				
	Length of Circuit: 180 cm				
	Connection Port(s): 15mm ID & 22mm OD				
TESTS PERFORMED ON THE	-The Leakage Test				
PRODUCT	-The Pull Test				
	-The Gauge Test				
	-The Routine Assembling And Packaging Process Controls				
APPLICABLE STANDARDS	Standard Number	Standard Name			
	TS EN ISO 5356-1:2015	Anaesthetic and respiratory equipment — Conical connectors — Part 1: Cones and sockets			
	TS EN ISO 11135:2014	Sterilization of health-care products — Ethylene oxide — Requirements for the development, validation and routine control of a sterilization process for medical devices			
	TS EN ISO 10993-1:2021	Biological evaluation of medical devices — Part 1: Evaluation and testing within a risk management process			
	TS EN ISO 10993-5:2010	Biological evaluation of medical devices — Part 5: Tests for in vitro cytotoxicity			
	TS EN ISO 10993-10:2014	Biological evaluation of medical devices — Part 10: Tests for irritation and skin sensitization			
	TS EN ISO 10993-12:2021	Biological evaluation of medical devices — Part 12: Sample preparation and reference materials			
	TS EN ISO 5362:2019	Anaesthetic reservoir bags			
	TS EN ISO 5367:2015	Anaesthetic and respiratory equipment - Breathing sets and connectors			
	ISO 13485:2016	Medical devices - Quality management systems - Requirements for regulatory			
		purposes			
	TS EN ISO 15223-1:2021	Medical devices — Symbols to be used with information to be supplied by the manufacturer — Part 1: General requirements			
	TS EN ISO 20417:2021	Medical devices - Information to be supplied by the manufacturer			
	TS EN ISO 14644-1:2016	Cleanrooms and associated controlled environments Part 1: Classification of air cleanliness			
	TS EN ISO 11607-1: 2020	Packaging for terminally sterilized medical devices - Part 1: Requirements for materials, sterile barrier systems and packaging systems			
	TS EN ISO 14971:2020	Medical devices - Application of risk management to medical devices			
	TS EN ISO 24971:2021	Medical devices — Guidance on the application of ISO 14971			
	TS EN ISO 10993-7:2010	Biological evaluation of medical devices part 7: Ethylene oxide sterilization residuals			
	TS EN ISO 10993-11: 2018	Biological evaluation of medical devices - Part 11: Tests for systemic toxicity			
	TS EN ISO 11737-1:2018	Sterilization of health care products — Microbiological methods — Part 1: Determination of a population of microorganisms on products			
	TS EN ISO 11737-2: 2020	Sterilization of health care products - Microbiological methods - Part 2: Tests of sterility performed in the definition, validation and maintenance of a sterilization process			
	TS EN 62366-1: 2015	Medical devices - Part 1: Application of usability engineering to medical devices			
STERILIZATION STATUS	Non-sterile				
CLEANING	Device assembled within ISC	0 8 Cleanroom.			
PRODUCT SHELF LIFE					
	5 years from the date of manufacturing. Expiration date and date of production are detailed on the product labelling.				
PACKAGING	Pouch: Polyethylene (PE) Box material: Craft Box dimention: 400 mm x 800 mm x 560 mm Quantity per box:				
STORAGE CONDITIONS	Temperature: -20°C to +55°C Humidity: 0% to 95% Luminosity: Keep away from direct sunlight				
TRANSPORTATION	Temperature: -20°C to +55°C				
CONDITIONS	Humidity: 0% to 95%				





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	Luminosity: Kee	Luminosity: Keep away from direct sunlight				
PRECAUTIONS	类	Keep away from sunlight	STERILEEO	Sterilized with Ethylene Oxide *for sterile products		
		Do not use if package is opened or damaged	€2195	CE Marking		
	2	Do not re-use	NON	Non-sterile *for non sterile products		
	PHT	Phthalate-free	LOT	Lot number		
		Consult instruction for use	REF	Catalog Number		
	LATEX	Latex-free		Expiry Date		
	erangen	Do not re sterilize *for sterile products	LATEX	Contains Latex *for products made with Latex		
	*55°C	Storage conditions				
	-20 °C	Country of manufacture – Date of manufacture				
	***	Manufacturer				
WASTE METHOD		Local regulations and/or hospital waste management procedures of the relevant country should be followed when disposing of the used products.				
NOTES	-	-				