

Product P/N	4222/701	Mod. 984A
Description	Eco Maxi Filter	Rev. 06



Eco Maxi Filter



PRODUCT	Inlet/Outlet Connectors: 22mm Male/15mm Female – 22mm Female/15mm Male ISO and
DESCRIPTION	Ø4.3mm ISO Luer Port.
	Approx. dimensions: 68.5mm diameter x 67.1mm height.
	Weight: 25g (approx.).
	Bidirectional Filter.
MANUFACTURER	GVS Filter Technology UK
NAME	NFC House
	Vickers Industrial Estate
	Mellishaw Lane, Morecambe
	Lancashire LA3 3EN - United Kingdom
	Information
	Tel. +44 (0) 1524 847600
	e-mail: gvsuk@gvs.com
INTENDED USE /	Filters protect the patient's airways effectively from exogenous microbial loads, thus reducing the
APPLICATION	risk of extrinsic colonisation and infection. Used to help reduce cross contamination between patient and machine. The luer port connector is used for monitoring respiratory and/or
	anaesthesia gases.
CLASS OF THE	Disposable medical device - Class Ila
PRODUCT	
	Rule 2 Annex IX 93/42 / EEC
	Rule 5 Annex VIII MDR 2017/745
MATERIALS	Filter media: Electrostatic Blended Synthetic Fiber
	Frame/Housing Polymer: Transparent Green Tinted Polypropylene (PP)
	Colour: Transparent Green
	Regulatory Documentation Required:
	- Biocompatibility according ISO 10993-1
	- ROHS
	- BSE/TSE
	- DEHP plasticizer Free and latex free
	- Aging
	- REACH
	- Conflict minerals
PRODUCT	Appearance/Visual
CHARACTERISTICS	As shown on drawing.
	Physical/Mechanical
	r nysica/mechanica



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	Approx. dimensions: 68.5mm diameter x 67.1mm height.
	Weight: 25g (approx.). Interfaces (ex: Input / Output connectors): 22mm Male/15mm Female – 22mm Female/15mm Male ISO and Ø4.3mm ISO Luer Port.
	Operating temperature Range: <i>N/A</i> Storage temperature Range: <i>5 °C to 40 °C</i> <i>Bidirectional Filter.</i>
	Biological Pyrogenicity: <0.3 EU/ml Biocompatibility to ISO10993 Category – Surface device Contact – Skin Contact Duration - <24hrs
	Functional Air Flow Rate: <i>301/min, 601/min, 901/min.</i>
	Filtration Efficiency: <i>NaCl Filter Efficiency</i> @ 30L/min using TSI 8130: Min. 98.5% (Ref.4222/700 REP: 0840/16 with factor of safety applied to Min.)
	Pressure Drop: Flow Resistance @ 30I/min in accordance with EN ISO 9360-1: Max. 110Pa Flow Resistance @ 60I/min in accordance with EN ISO 9360-1: Max. 231Pa Flow Resistance @ 90I/min in accordance with EN ISO 9360-1: Max. 374Pa (REP:0708/16 with 10% of safety margin added to Max.)
	Internal Volume: 28ml (approx.)
	Operating Lifetime: Refer to Instructions for Use.
	Shelf Lifetime: 5 years from the date of manufacture.
	Bacterial Filtration Efficiency in accordance with ASTM F2101-07: Min. 99.999% (Staphylococcus aureus @ 30L /minute) REP: EXT486704B
	Viral Filtration Efficiency in accordance with ASTM F2101-07: Min. 99.999% (Bacteriophage @ 30L/ minute) REP: EXT486705B.1
	Gas leakage in accordance with EN9360: Max.0.0 ml/min (REP:1309/17)
	Cleanliness Device assembled within Class 8 Cleanroom.
	Testing Leak test at 3PSI.
INSTRUCTIONS / WARNINGS	Multi-language IFU available.
PRODUCT SHELF LIFE	5 years from the date of manufacture.



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	Expiration date and date of manufacture are detailed on the product labelling.
STERILIZATION	Sterile version of product available (Ethylene oxide - Max 55°C)
APPLICABLE STANDARDS AND REGULATIONS	Product Certification required: - CE mark - FDA
	Applicable Standards and Technical Regulations: Biological evaluation of Medical Devices - Part 1: Evaluation and Testing - ISO 10993-1.
	Respiratory protective devices - Method for test - Part 7: Determination of particle filter penetration - BS EN 13274-7.
	Medical devices- Application of risk management to medical devices - BS EN ISO 14971.
	Medical devices – symbols to be used with medical device labels, labelling and information to be supplied - Part1: General requirements - ISO 15223-1.
	Breathing system filters for anaesthetic and respiratory use — Part 1: Salt test method to assess filtration performance - ISO 23328-1.
	Anaesthetic and respiratory equipment – conical connectors – part 1: Cones and sockets – ISO 5356-1.
	Sterilization of health care products – Ethylene oxide sterilization – ISO 11135-1.
	Sterilization of medical devices – Microbiological Methods – Part 1: Estimation of population of microorganisms on products – ISO 11737-1.
PACKAGING AND LABELING	 Number of pcs per bag is determined by the sales order. The first barcode label is applied to the outside of the bags. The second barcode label is applied onto the outside of the box. Each bag is labelled with the following traceability information: Quantity Product description Product Date Lot Number (OL and 5-digit batch number to trace back to raw materials used) Operator Code Different lots in one box are separately closed and separately labelled. Bulk products will be packed in double PE bags.
CERTIFICATE OF COMPLIANCE	With each shipment, GVS UK Customer Service will send the CofC to the Customer, based on the lot numbers and date of manufacture. Conformity declaration is printed on every invoice and Certificate is according to UNI EN 10204 type 2.1. The Quality management system is in compliance with ISO 13485.
DRAWING	The attached drawing is part of this product specification and must not be duplicated or made accessible to a third party without written permission from GVS Filter Technology UK Ltd.



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	22 Female / 15 Male Connector in accordance with EN ISO 5356 22 Male / 15 Female Connector in accordance with EN ISO 5356		67.1 67.1
ACCEPTABLE QUALITY LEVEL	AQL: 0.65 with sampling Plan: ISO2859.		
VISUAL REQUIREMENTS	Visual acceptance requirements apply when inspected under be Magnification: Unaided eye at a distance of approximately 35-40cm. Light type: Lighting level must be reasonable for visual detection. Timings: Maximum inspection period per item is 25 seconds. For detailed defect list, refer to product control plan.		
	Acceptance Requirement	AQL	Sampling Plan
	1 Black particle contamination	0.65	
	2 Damaged/broken item	0.65	ISO 2859 Part 1
	3 Blocked connector/luer	0.65	General Inspection
	4 Weld marks	0.65	



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	5	5 Short fill moulding	0.65	
	e	5 Rough surface or edges	0.65	
	7	Pronounced injection gate	0.65	
	8	B Deformation/distortion	0.65	
	9	P Crack	0.65	
	1	0 Oil/grease	0.65	
	1	1 Wrong colour	0.65	
	1	2 Weld fault	0.65	
GENERAL SAFETY AND PERFORMANCE	Special characteristic: Product characteristic which can affect safety or compliance with regulations, fit, function, performance or subsequent processing of product. Special Characteristic # 01:			
REQUIREMENTS	Flow	Resistance @ 30L/min in accordance with EN ISO 9360-1,		
	Flow	Resistance @ 60L/min in accordance with EN ISO 9360-1,		
	Flow	Resistance @ 90L/min in accordance with EN ISO 9360-1.		
Special Characteristic # 02: NaCl Filter Efficiency @ 30L/min using TSI 8130 in acc with EN 13274-7.		TSI 8130 in accor	dance	
	Spec	ial Characteristic # 03: Bacterial Filtration Efficiency in accord	ance with ASTM F2	2101-07,
	Viral	Filtration Efficiency in accordance with ASTM F2101-07.		
	Spec	ial Characteristic # 04: Conical connectors compliant in accord	ance with EN5356	i.
	Special Characteristic # 05: Gas Leakage compliant in accordance with EN9360.			
This material specific	cation	describes the properties of product above indicated. This d	ocument contains	s general

This material specification describes the properties of product above indicated. This document contains general requirements, material description, drawing references, defect specification, biological material requirements.

REVISIONS AND APPROVALS:

DATE	REV.	REASON FOR CHANGE	ISSUED AND CONTROLLED BY: (NAME/FUNCTION/SIGNATURE)	APPROVED BY: (NAME/FUNCTION/SIGNATURE)
03/08/2021	4	Functional characteristics updated.	Kinga Gawdzik – Engineering Support Technician	Andrew Pearce – Quality Manager



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CUSTOMER APPROVAL:		
le accept this material specification as a part of the agreed terms of delivery.		
ompany Name:		
pproved by:		
NAME/FUNCTION		
SIGNATURE		
DATE		
COMPANY STAMP		

Please send back this document signed for approval. If we will not receive this specification signed, we consider the first order placed as implicit approval.