

PRODUCT SPECIFICATION

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|-------------|-------------------------------------|----------------------|
| Product P/N | 2800/23 | Mod. 984A Rev. 06 |
| Description | Spiroguard with Integral Mouthpiece | |

2800/23

***Spiroguard
with Integral
Mouthpiece***



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| PRODUCT DESCRIPTION | <p>Inlet/Outlet Connectors: - OD 48.36mm / ID 45.35mm - Machine Side; - Integral Mouthpiece - Patient Side. Approx. dimensions: 96.8mm diameter x 79.2mm height. Weight: 37gm (approx.). Bidirectional Filter.</p> |
| MANUFACTURER NAME | <p>GVS Filter Technology UK NFC House Vickers Industrial Estate Mellishaw Lane, Morecambe Lancashire LA3 3EN - United Kingdom</p> <p>Information Tel. +44 (0) 1524 847600 e-mail: gvsuk@gvs.com</p> |
| INTENDED USE / APPLICATION | <p>Spirometer pre filters are used in pulmonary function tests for trapping bacteria and viruses and other particulates, reducing the risk of cross-contamination between the patient and the machine.</p> |
| CLASS OF THE PRODUCT | <p>Disposable medical device - Class IIa Rule 2 Annex IX 93/42 / EEC Rule 5 Annex VIII MDR 2017/745</p> |
| MATERIALS | <p>Filter media: <i>Electrostatic Blended Synthetic Fiber</i> Frame/Housing Polymer: <i>White Polypropylene Homopolymer (PP)</i> Colour: <i>White</i></p> <p>Regulatory Documentation Required:</p> <ul style="list-style-type: none"> - Biocompatibility according ISO 10993-1 - ROHS - BSE/TSE - DEHP plasticizer Free and latex free - Aging - REACH - Conflict minerals |

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| PRODUCT CHARACTERISTICS | <p>Appearance/Visual As shown on drawing.</p> <p>Physical/Mechanical Approx. dimensions: 96.8mm diameter x 79.2mm height. Weight: 37gm (approx.). Interfaces (ex: Input / Output connectors): OD 48.36mm / ID 45.35mm - Machine Side; Integral Mouthpiece - Patient Side. Operating temperature Range: N/A Storage temperature Range: 5 °C to 40 °C Bidirectional Filter, Integral Mouthpiece – Patient Side.</p> <p>Biological Pyrogenicity: <0.3 EU/ml Biocompatibility to ISO10993 Category – Surface device Contact – Oral cavity Contact Duration - <24hrs</p> <p>Functional Air Flow Rate: 30l/min, 60l/min, 90l/min. Filtration Efficiency: Filter Efficiency @ 30L/min using TSI 8130: Min. 99% (REP: 1433/17 with factor of safety applied to Min.) Pressure Drop: Flow Resistance @ 30l/min in accordance with EN ISO 9360-1: Max.33Pa Flow Resistance @ 60l/min in accordance with EN ISO 9360-1: Max.68.2Pa Flow Resistance @ 90l/min in accordance with EN ISO 9360-1: Max.112.2Pa (REP:1430/17 with 10% of safety margin added to Max.) Internal Volume: 68ml (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: EXT607770. Viral Filtration Efficiency in accordance with ASTM F2101-07: Min. 99.999% (Bacteriophage @ 30L/ minute) REP: EXT620332.</p> <p>Cleanliness Device assembled within Class 8 Cleanroom.</p> <p>Testing Torque test @ 10Nm. (REP: 1431/17)</p> |
|-------------------------|--|

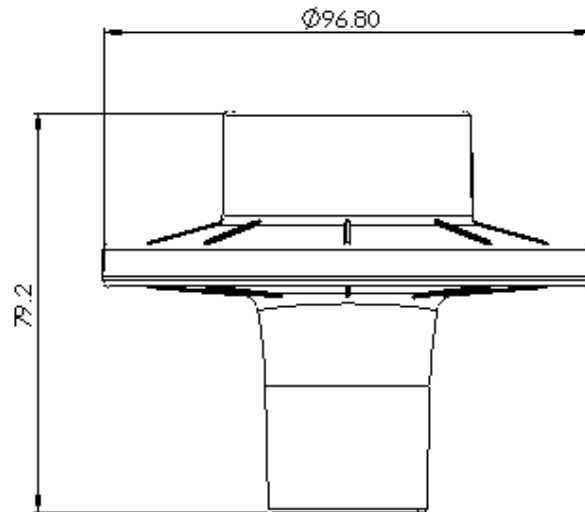
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| INSTRUCTIONS / WARNINGS | <i>Multi-language IFU available.</i> |
| PRODUCT SHELF LIFE | <p><i>5 years from the date of manufacture.</i></p> <p><i>Expiration date and date of manufacture are detailed on the product labelling.</i></p> |
| STERILIZATION | <i>N/A</i> |
| APPLICABLE STANDARDS AND REGULATIONS | <p>Product Certification required:</p> <ul style="list-style-type: none"> - CE mark - FDA <p>Applicable Standards and Technical Regulations:</p> <p><i>Biological evaluation of Medical Devices - Part 1: Evaluation and Testing - ISO 10993-1.</i></p> <p><i>Respiratory protective devices - Method for test - Part 7: Determination of particle filter penetration - BS EN 13274-7.</i></p> <p><i>Medical devices- Application of risk management to medical devices - BS EN ISO 14971.</i></p> <p><i>Medical devices – symbols to be used with medical device labels, labelling and information to be supplied - Part1: General requirements - ISO 15223-1.</i></p> |
| PACKAGING AND LABELING | <p><i>Number of pcs per bag is determined by the sales order.</i></p> <p><i>The first barcode label is applied to the outside of the bags.</i></p> <p><i>The second barcode label is applied onto the outside of the box.</i></p> <p><i>Each bag is labelled with the following traceability information:</i></p> <ul style="list-style-type: none"> ✓ Quantity ✓ Product description ✓ Product Date ✓ Lot Number (OL and 5-digit batch number to trace back to raw materials used) ✓ Operator Code <p><i>Different lots in one box are separately closed and separately labelled.</i></p> <p><i>Bulk products will be packed in double PE bags.</i></p> |
| CERTIFICATE OF COMPLIANCE | <p><i>With each shipment, GVS UK Customer Service will send the CofC to the Customer, based on the lot numbers and date of manufacture.</i></p> <p><i>Conformity declaration is printed on every invoice and Certificate is according to UNI EN 10204 type 2.1.</i></p> <p><i>The Quality management system is in compliance with ISO 9001, ISO 13485.</i></p> |
| DRAWING | <i>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 Lt</i> |

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Approximate dimensions for reference only

ACCEPTABLE QUALITY LEVEL

AQL: 0.65 with sampling Plan: ISO2859.

VISUAL REQUIREMENTS

Visual acceptance requirements apply when inspected under below conditions:

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 | ISO 2859 Part 1 General Inspection Level 1 |
| 2 | Damaged/broken item | 0.65 | |
| 3 | Blocked connector/luer | 0.65 | |
| 4 | Weld marks | 0.65 | |
| 5 | Short fill moulding | 0.65 | |
| 6 | Rough surface or edges | 0.65 | |
| 7 | Pronounced injection gate | 0.65 | |
| 8 | Deformation/distortion | 0.65 | |

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|---|--|--------------|------|--|--|
| | 9 | Crack | 0.65 | | |
| | 10 | Oil/grease | 0.65 | | |
| | 11 | Wrong colour | 0.65 | | |
| | 12 | Weld fault | 0.65 | | |
| GENERAL SAFETY AND PERFORMANCE REQUIREMENTS | Special characteristic: <i>Product characteristic which can affect safety or compliance with regulations, fit, function, performance or subsequent processing of product.</i> | | | | |
| | Special Characteristic # 01: | | | | |
| | <i>Flow Resistance @ 30L/min in accordance with EN ISO 9360-1</i> | | | | |
| | <i>Flow Resistance @ 60L/min in accordance with EN ISO 9360-1</i> | | | | |
| | <i>Flow Resistance @ 90L/min in accordance with EN ISO 9360-1</i> | | | | |
| | Special Characteristic # 02: <i>Filter Efficiency @ 30L/min using TSI 8130 in accordance with EN 13274-7.</i> | | | | |
| Special Characteristic # 03: <i>Bacterial Filtration Efficiency in accordance with ASTM F2101-07</i> | | | | | |
| <i>Viral Filtration Efficiency in accordance with ASTM F2101-07</i> | | | | | |
| 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) |
|------------|------|----------------------------|---|--|
| 11/08/2021 | 3 | Filter Efficiency updated. | Kinga Gawdzik – Engineering Support Technician  | Andrew Pearce – Quality Manager  |

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CUSTOMER APPROVAL:

We accept this material specification as a part of the agreed terms of delivery.

Company Name:

Approved 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.