

Data Sheet

AseptiSure® HS

High Temperature Resistant Hydrophilic Polyethersulfone (PES) Membrane Cartridge Filters

Biopharmaceutical processing requires sterilizing grade microfiltration at multiple stages to meet specific process requirements.

Processes managers are continuously looking for microfiltration solutions to upstream, downstream, intermediate processes and final biological preparations. Since bio manufacturing is a multi stage process and bio molecules by nature are extremely sensitive, they are looking for:

- Minimizing protein losses due to adsorption to improve over all product yields
- Minimizing filter extracts which add up due to multiple points of use in a process
- High throughputs to achieve process economy
- Absolute retentions for higher sterility assurance

mdi *AseptiSure® HS* Polyethersulfone (PES) Membrane Cartridge filters are high temperature resistant filtration devices. These are designed to withstand high pressure differential at high temperature steam sterilization up to 135°C.

AseptiSure® HS cartridge filters with polyethersulfone membrane serial layers offer enhanced throughputs, thus ensuring better economics. These are validated for key performance parameters such as retention efficiency, chemical compatibility, extractables, heat stability and flow rates.



AseptiSure® HS

High Temperature Resistant PES Membrane Cartridge Filters

Datasheet

mdi AseptiSure® HS cartridge filters are deeply validated for use in Biopharmaceutical applications. These filters are manufactured in class 10,000 clean rooms and ISO 9001 certified facilities.

Key Features

- Low protein binding
- High throughputs
- Long service life
- Pre-flushed to minimize particulate release after installation
- Non-toxic material of construction
- Absolute retention
- 100% integrity tested
- High flow rates
- Bioburden maintained below 1000 cfu/device
- Endotoxin level certified to be <0.25 EU/ml
- Unique identification number is laser etched on each filter
- Individual certificate of quality for each device
- Sterilizable by Autoclaving/Steaming in place (SIP)

Applications

- Sterile filtration of:
Proteinaceous liquids where minimum protein loss is desired, such as sera, culture soups and recombinant proteins, antibodies etc
- Cell culture media
- Buffers
- Small volume parenterals
- Large volume parenterals

Validation Services

The regulatory requirements emphasize on the need to validate the efficacy of the filter with drug product under simulated worst-case conditions of use.

mdi provides validation services supported by customized validation protocols and world class test facilities to assist you in filter validations with your specific drug product.

mdi quality management system emphasizes on quality by design rather than by end product testing. Robust processes are developed for product manufacturing and are continuously monitored to ensure that the products meet their predetermined specifications and lot to lot reproducibility is ensured.

Certificate of Quality

Each cartridge filter is accompanied by individual certificate of quality to ensure traceable documentation at user's end.

It certifies the product compliance to various regulatory as well as user requirements.

Validated for Microbial Retention

Integrity test data have been correlated to actual microbial retention with *B.diminuta* (ATCC 19146) as per ASTM F838 to establish acceptable integrity test values.

Samples from each lot are subjected to microbial challenge test before final lot release.

100% Integrity Tested

Each *AsepsiSure*® HS is tested for integrity to comply with validated Acceptable Integrity Test Specifications.

Flow Rate

Each lot is tested for clean water flow rates to ensure that flow rates are within the specifications.

Adsorption

AsepsiSure® HS filters are validated for low protein binding to ensure minimal active ingredient losses when used for filtration of high value proteins.

Pressure, Temperature Endurance

AsepsiSure® HS filters are validated to endure high operating pressure and temperature conditions which may be encountered during use.

Extractables

Extractables/leachables from *AsepsiSure*® HS filters, used at various stages of a biopharmaceutical manufacturing process, will add on and may impact the impurity profile of the desired product.

AsepsiSure® HS filters are validated to exhibit low extractables under harsh extraction conditions.

Bioburden Testing

AsepsiSure® HS bioburden is tested as per ISO 11737-1 and assured to be <1000 cfu/device.

Endotoxin Testing

Aqueous extracts exhibit < 0.25 EU/ml as established by Limulus Amebocyte Lysate (LAL) Test as per USP <85>.

Total Traceability

AsepsiSure® HS filters come with completely traceable lot numbers and unique identification number to facilitate easy and fast retrieval of manufacturing and quality control data associated with each filter.

These unique lot and identification numbers are laser etched on each filter device and also printed on the labels of the box in which individual filter is packed.

Packaging Integrity

AsepsiSure® HS filters are packed in bags to ensure package integrity during transit as well as to prevent particulate contamination while transferring to clean room process areas.

Other Regulatory Compliance

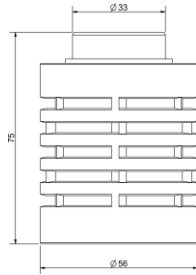
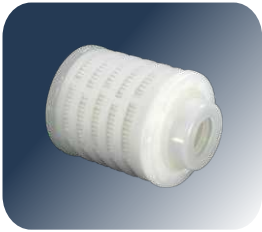
- Complies with USFDA 21 CFR 210.3(b)(6) for fiber release
- Complies with USFDA 21 CFR 177.1520 for fractional dissolution
- Materials of construction tested for toxicity as per Biological Reactivity Tests, In-vivo, USP <88> for class VI Plastics
- Complete filter devices tested for cytotoxicity as per Biological Reactivity Tests, In-vitro, USP <87>

Adapters and Dimensions

Datasheet

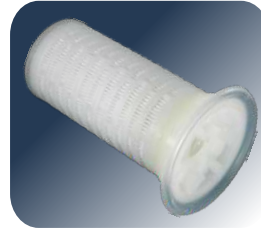
2.5" Mini Cartridge Filters

4463 Adapter (E0)

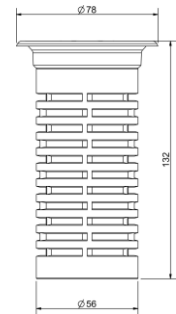


Total Length : 75 mm
Diameter : 56 mm

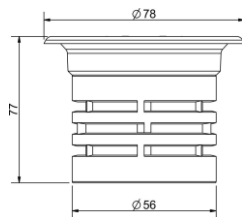
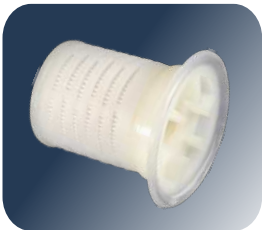
Seal-K Adapter (G0)



Total Length : 132 mm
Diameter : 56 mm

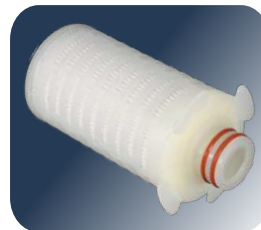


Seal-K Adapter (G0)

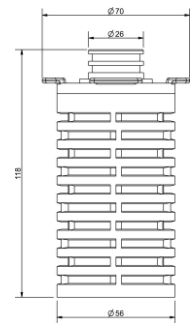


Total Length : 77 mm
Diameter : 56 mm

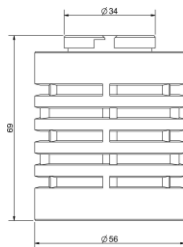
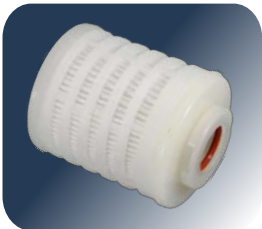
4440 Adapter (U0)



Total Length : 118 mm
Diameter : 56 mm



4463B Adapter (H0)

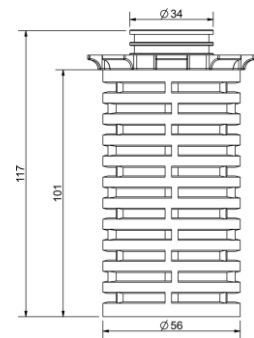


Total Length : 69 mm
Diameter : 56 mm

Seal-O Adapter (F0)

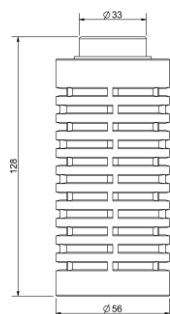
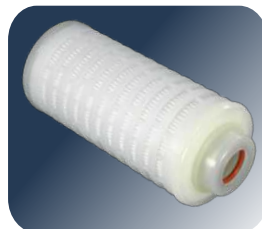


Total Length : 117 mm
Diameter : 56 mm



5" Mini Cartridge Filters

4463 Adapter (E0)

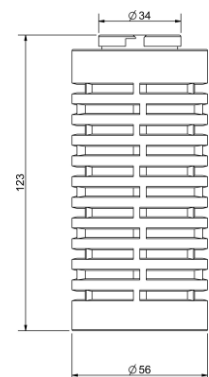


Total Length : 128 mm
Diameter : 56 mm

4463B Adapter (H0)

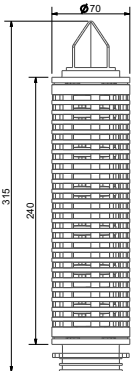


Total Length : 123 mm
Diameter : 56 mm



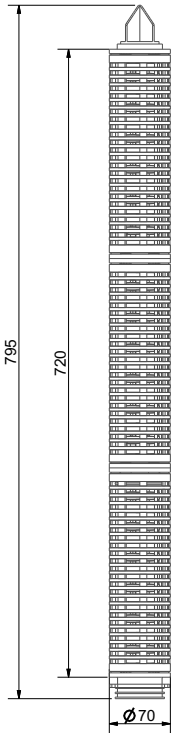
Standard Cartridge Filters

10" Cartridge Filter- 7P Adapter with Fin (A0)



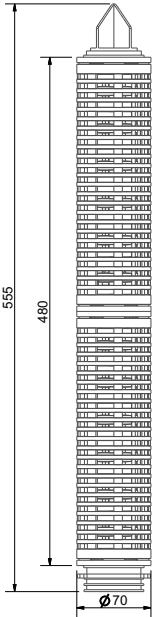
Total Length : 315 mm
Diameter : 70 mm

30" Cartridge Filter- 7P Adapter with Fin (A0)



Total Length : 795 mm
Diameter : 70 mm

20" Cartridge Filter- 7P Adapter with Fin (A0)



Total Length : 555 mm
Diameter : 70 mm

Adapter and Elastomers Availability Chart

Datasheet

Mini Cartridge Filters		
Adapters	2.5"	5"
4463	√	√
4463B	√	√
4440	√	√
Seal-K	√	√
Seal-O	X	√
Seal-M	√	√

Mini Cartridge Filters	
Adapters	Elastomer
	Silicone
4463	√
4463B	√
4440	√
Seal-K	X
Seal-O	√
Seal-M	√

Standard Cartridge Filters				
Adapters	5"	10"	20"	30"
7P	√	√	√	√
7P without Fin	√	√	√	√
28 with Fin	X	√	√	√
'O'	X	√	√	√

Standard Cartridge Filters				
Adapters	Elastomers			
	Silicone	Viton	EPDM	FEP Encapsulated Viton
7P	√	√	√	√
7P without Fin	√	√	√	√
28 with Fin	√	√	√	X
'O'	√	√	√	X

Linear Upscaling from Pilot Scale to Production Process

Datasheet

Scientists are concerned about filter fluid interaction impacting the stability, purity, strength etc. of the drug product, and they take a keen interest in filter selection at the formulation development stage itself. Although preliminary compatibility data support initial filter selection, for stability studies detailed filter validations are required to provide enough documented evidence to justify specific filter use.

A critical requirement that needs to be addressed at this stage is of scalability from pilot scale to full scale production processes.

mdi offers a wide range of *AseptiSure® HS* filters to provide linear scale up from lab scale to production process. While scaling up the process, the appropriate size filter can be selected by increasing the effective filtration area of filter proportionate to the process fluid volumes.

All Materials of construction as well as manufacturing process are identical for all filter devices starting from 1000 cm² to 18000cm² hence process scaling can be facilitated without triggering additional validation studies for given process conditions. **mdi** provides complete documentation for each of the *AseptiSure® HS* filters there by reducing the additional validation cost and time.



AseptiSure® HS, 2.5"
EFA: 1000 cm²



AseptiSure® HS, 5"
EFA: 2000 cm²



AseptiSure® HS, 5" Large
EFA: 3000 cm²



AseptiSure® HS, 10"
EFA: 6000 cm²



AseptiSure® HS, 20"
EFA: 12000 cm²



AseptiSure® HS, 30"
EFA: 18000 cm²

*EFA: Effective Filtration Area

Specifications

Mini Cartridge Filters

Datasheet

Construction

Membrane	Hydrophilic PES	
Support Layers	Polyester	
Plastic Parts	Polypropylene	
O rings	Silicone	
Final Filter Pore Size	0.2µm	0.45µm
Pre-Filter Pore Size	0.8 µm, 0.65 µm, 0.45 µm	0.65 µm, 0.8 µm

Integrity Testing / Retention

Pore Size	0.2µm	0.45µm
Bubble Point	≥ 50psi (3.52Kg/cm ²) with Water	≥ 30psi (2.11Kg/cm ²) with Water
Microbial Retention	LRV >7 for <i>Brevundimonas diminuta</i> (ATCC 19146) per cm ²	LRV >7 for <i>Serratia marcescens</i> (ATCC 14756) per cm ²

Size

Size	2.5"	5"
Effective Filtration Area (Nominal)	1000cm ²	2000cm ²

Operational

Max. Operating Temperature	80 °C @ < 30 psi (2 Kg/cm ²)
Max. Differential Pressure	50 psi (3.5 Kg/cm ²) @ 25 °C
Reverse Pressure	< 0.7 Kg/cm ² (10 psi) @ 25 °C
Sterilization	Autoclavable/In-line steam sterilizable at 135 ° C for 30 minutes, 5 cycles

Assurance

Toxicity	Passes Biological Reactivity tests, In Vivo, as per USP <88> for Class VI plastics
Cytotoxicity	Passes Biological Reactivity tests, In Vitro, USP <87> for cytotoxicity
Bacterial Endotoxin	Aqueous extracts exhibit < 0.25 EU/ml as established by Limulus Amebocyte Lysate (LAL) Test as per USP <85>
Bioburden	Bioburden level is < 1000 cfu/filter device as per ISO 11737-1
Particle Shedding	The filtrate complies with USP <788> test for particulate matter in injections
Non Fiber Releasing	Passes test as per USP and comply with USFDA 21 CFR Part 210.3(b)(6) for fiber release
TOC and Conductivity	Meets the WFI requirements of USP for TOC <643> and Conductivity <645> after a 3 liter WFI flush
Extractables with WFI	Passes NVR test as per USP <661>
pH Compatibility	Compatible with pH range of 1-10
Indirect Food Additives	All Polypropylene components meet the FDA Indirect Food Additive requirements cited in 21 CFR 177.1520
Oxidizable Substances	Passes test as per USP <1231>
Quality Management System	ISO-9001 Certified
USFDA	DMF No. 015554

Specifications

Standard Cartridge Filters

Datasheet

Construction

Membrane	Hydrophilic PES		
Support Layers	Polyester		
Plastic Parts	Polypropylene		
O rings	Silicone		
	Viton		
	EPDM		
	FEP Encapsulated Viton		
Final Filter Pore Size	0.2µm	0.45µm	
Pre-Filter Pore Size	0.8 µm, 0.65 µm, 0.45 µm	0.65 µm, 0.8 µm	

Integrity Testing / Retention

Pore Size	0.2µm	0.45µm	
Bubble Point	≥ 50psi (3.52Kg/cm ²) with Water	≥ 30psi (2.11Kg/cm ²) with Water	
Air Diffusion Flow (10" Filter)	≤ 30 ml/min @ 37 psi (2.6 Kg/cm ²) with Water	≤ 35 ml/min @ 22 psi (1.54 Kg/cm ²) with Water	
Microbial Retention	LRV >7 for <i>Brevundimonas diminuta</i> (ATCC 19146) per cm ²	LRV >7 for <i>Serratia marcescens</i> (ATCC 14756) per cm ²	

Size

Size	5"	10"	20"	30"
Effective Filtration Area (Nominal)	3000cm ²	6000cm ²	12000cm ²	18000cm ²

Operational

Max. Operating Temperature	80 °C @ < 30 psi (2 Kg/cm ²)		
Max. Differential Pressure	50 psi (3.5 Kg/cm ²) @ 25 °C		
Reverse Pressure	< 0.7 Kg/cm ² (10 psi) @ 25 °C		
Sterilization	Autoclavable/In-line steam sterilizable at 135 ° C for 30 minutes, 5 cycles		

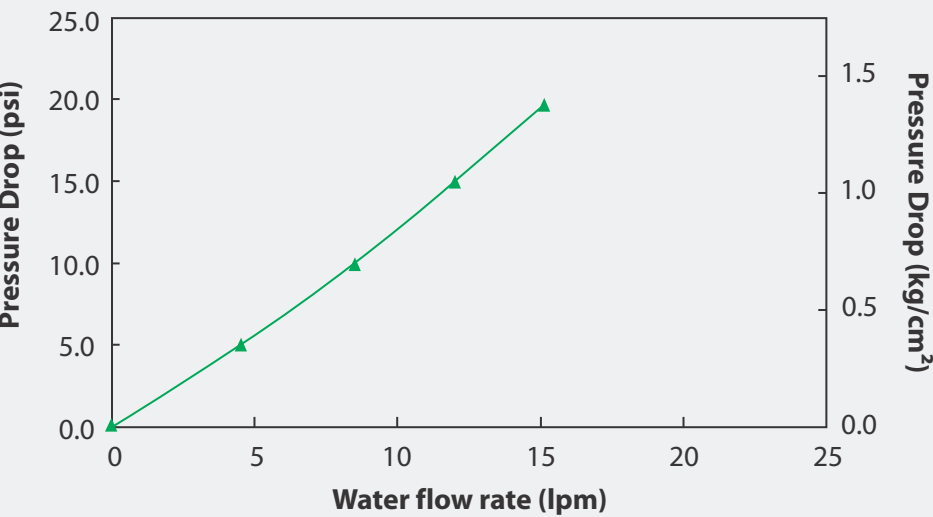
Assurance

Toxicity	Passes Biological Reactivity tests, In Vivo, as per USP <88> for Class VI plastics		
Cytotoxicity	Passes Biological Reactivity tests, In Vitro, USP <87> for cytotoxicity		
Bacterial Endotoxin	Aqueous extracts exhibit < 0.25 EU/ml as established by Limulus Amebocyte Lysate (LAL) Test as per USP <85>		
Bioburden	Bioburden level is < 1000 cfu/filter device as per ISO 11737-1		
Particle Shedding	The filtrate complies with USP <788> test for particulate matter in injections		
Non Fiber Releasing	Passes test as per USP and comply with USFDA 21 CFR Part 210.3(b)(6) for fiber release		
TOC and Conductivity	Meets the WFI requirements of USP for TOC <643> and Conductivity <645> after a 3 liter WFI flush		
Extractables with WFI	Passes NVR test as per USP <661>		
pH Compatibility	Compatible with pH range of 1-10		
Indirect Food Additives	All Polypropylene components meet the FDA Indirect Food Additive requirements cited in 21 CFR 177.1520		
Oxidizable Substances	Passes test as per USP <1231>		
Quality Management System	ISO-9001 Certified		
USFDA	DMF No. 015554		

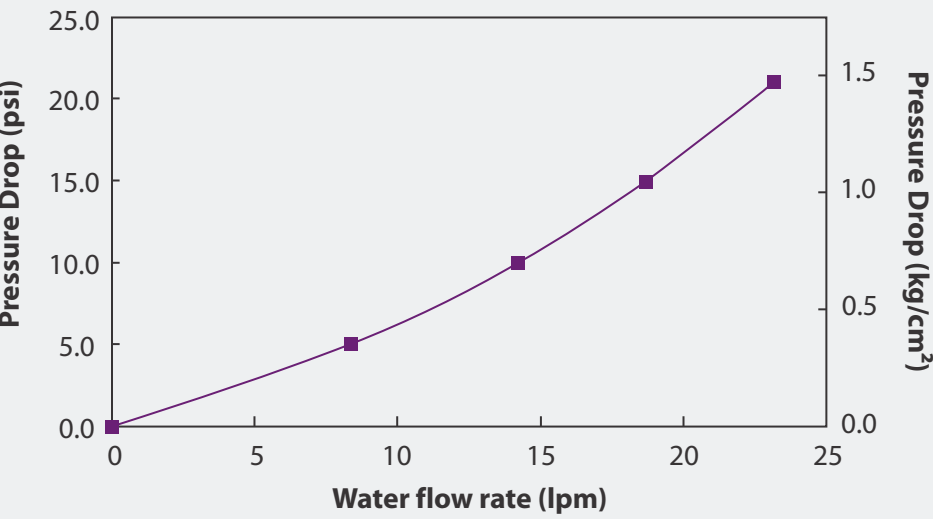
Typical Water Flow Rates

Mini Cartridge Filters

0.2µm AseptiSure® HS, 2.5" Mini Cartridge Filters



0.2µm septiSure® HS, 5" Mini Cartridge Filters

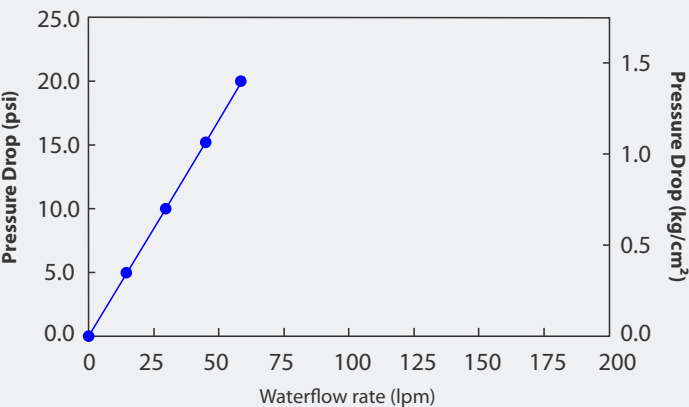


Typical Water Flow Rates

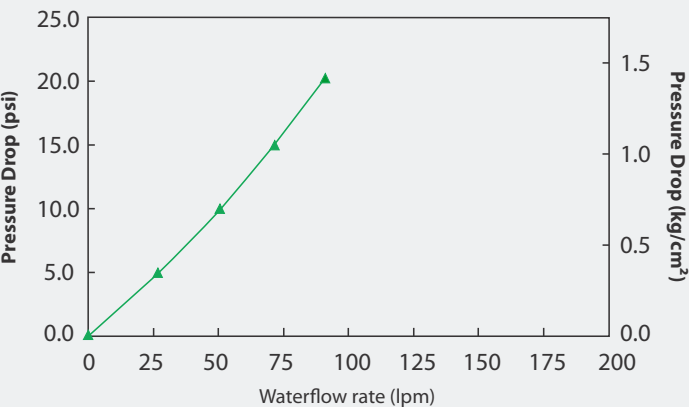
Standard Cartridge Filters

Datasheet

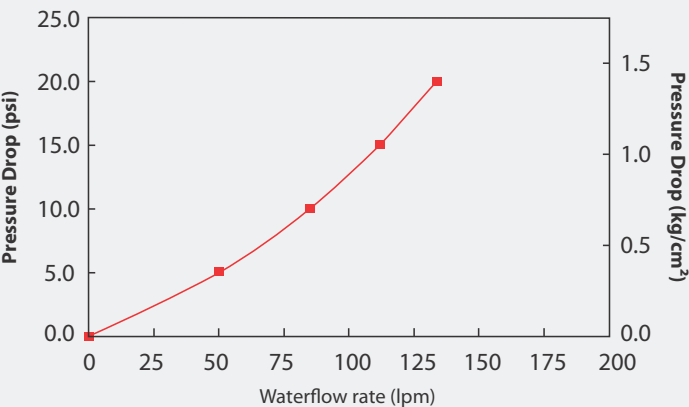
0.2µm AseptiSure® HS, 5" Standard Cartridge Filters



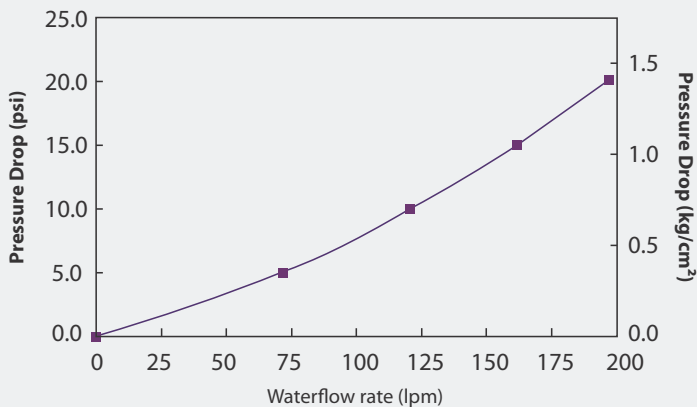
0.2µm AseptiSure® HS, 10" Standard Cartridge Filters



0.2µm AseptiSure® HS, 20" Standard Cartridge Filters



0.2µm AseptiSure® HS, 30" Standard Cartridge Filters



Ordering Information

Datasheet

AseptiSure® HS PES Membrane Mini Cartridge Filter

Type		Size		Pore Size		Adapter		Elastomer		Sterility		Pack Size	
	Code		Code		Code		Code		Code		Code		Code
AseptiSure® HS (0.8µm Upstream)	CPH5	2.5"	50	0.2µm	01	4463	E0	Silicone	SS	Non Sterile	1	1	01
AseptiSure® HS (0.65µm Upstream)	CPH3	5"	53	0.45µm*	02	4463B	H0						
AseptiSure® HS (0.45µm Upstream)	CPHX					4440	U0						
						Seal-K	G0**						
						Seal-O	F0***						
						Seal-M	J0						

Example:

CPHX	50	01	E0	SS	1	01
------	----	----	----	----	---	----

*0.45µm cartridge filters are available with 0.65µm or 0.8µm upstream layer only

**G0 adapter code is not available with any elastomer. Please mention XX in place of elastomer code while ordering

*** Adapter code F0 is available only in 5" cartridge filters.

AseptiSure® HS PES Membrane Standard Cartridge Filter

Type		Size		Pore Size		Adapter		Elastomer		Sterility		Pack Size	
	Code		Code		Code		Code		Code		Code		Code
AseptiSure® HS (0.8µm Upstream)	CPH5	5"	53	0.2µm	01	7P	A0	Silicone	SS	Non Sterile	1	1	01
AseptiSure® HS (0.65µm Upstream)	CPH3	10"	54	0.45µm**	02	7P without fin	A1	EPDM	SE				
AseptiSure® HS (0.45µm Upstream)	CPHX	20"	55			28 with fin	C0	Viton	SV				
		30"	56			'O'	D0	FEP Encapsulated Viton	FV***				

Example:

CPHX	54	01	A1	SS	1	01
------	----	----	----	----	---	----

*Size 5" is available in Adapter Code A0 (7P) and A1 (7P without fin) only

**0.45µm Cartridge filters are available with 0.65µm or 0.8µm upstream layer only

***FV is available in adapter code A0 (7P) and A1 (7P without fin) only

Advanced Microdevices Pvt. Ltd.

20-21, Industrial Area, Ambala Cantt-133 006, INDIA

Tel : +91-171-2699290, 2699471

E-mail : info@mdimembrane.com

Website : www.mdimembrane.com



Distribuidor Exclusivo España

Avenida Cerdanyola, 92-94 Local B4

08173 - Sant Cugat del Vallès - Barcelona

Tel: 935921554 - Email: comercial@gesfilter.com

www.gesfilter.com