

AseptiCap WS Hydrophilic PVDF Membrane Capsule Filters

mdi AseptiCap WS are low protein binding hydrophilic PVDF membrane capsule filters offering serial filtration incorporating a large pore size upstream membrane to protect the downstream membrane for enhanced throughput.

These capsule filters are validated to meet compendia and regulatory requirements and are well characterized. They meet key process requirements such as absolute retention efficiency, extremely low extractables, high throughputs, wide chemical compatibility and other important characteristics.

Key features

- Absolute retention
- > 100% integrity tested
- Low protein binding
- Low extractables
- Very low hold up volume in filters

Applications

Sterile Filtration of

- > Antibodies
- Protein Solutions
- Buffers
- Vaccine concentrates
- Small Volume Parenterals

Microbially Validated as per ASTM F 838-05 Complies with USFDA 21 CFR 210.3(b)(6) Meets and Exceeds USFDA 21 CFR 177.1520





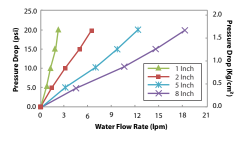
Specifications

Pore Size 0.2 μm and 0.45 μm

Materials of Construction

Membrane	Hydrophilic PVDF
Support Layer	Polyester
Plastic Components	Polypropylene

Typical Water Flow Rates, 0.2 μm



Microbial Retention

0.2 \mum: LRV >7 for *B. diminuta* (ATCC 19146) per cm²

0.45 \mum: LRV >7 for *S. marcescens* (ATCC 14756) per cm²

Maximum Operating Temperature

80 °C @ <u><</u> 30 psi (2 Kg/cm²)

Maximum Differential Pressure 60 psi (4 Kg/cm²) @ 30 °C

Bubble Point (with water)

0.2 μm: ≥ 50psi (3.51Kg/cm²) **0.45 μm**: ≥ 30 psi (2.11 Kg/cm²)

Sterilization

By Autoclave: Autoclavable at 125°C for 30 minutes, 2 Cycles. Can not be in-line steam sterilized

By Gas: Sterilization by Ethylene Oxide

Toxicity

Passes Bioreactivity test, 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>

Fiber Release

Passes test as per USP and comply with USFDA 21 CFR Part 210.3(b)(6) for fiber release

Particle Release:

The filtrate complies with USP <788> test for particulate matter in injections

TOC and Conductivity

Meets the WFI requirements of USP for TOC <643> and Conductivity <645> after a 3 liter flush

Extractables with WFI

Passes NVR test as per USP <661>

Oxidizable Substances

Passes test as per USP <1231>

Туре		Size		Pore Size		Inlet /Outlet		Х	x	Sterility		Pack Size	
	Code	Length and EFA	Code		Code		Code				Code		Code
AseptiCap WS (0.45 μm Upstream) D	DWSX	1" (250 cm ²)	51	0.2 µm	01	1⁄4″ SHB	A			Non-Sterile	1	1	01
		2" (500 cm ²)	52	0.45 µm	02	½" Hose Barb	D			EO Sterile	2		
AseptiCap WS (0.8 μm Upstream)*	DWS5	5" (1000 cm ²)	53			1½" Sanitary Flange	E					-	
	00055	8" (2000 cm ²)	57			³ ⁄ ₄ " Sanitary Flange	S	$*0.8\mu m$ upstream is available with 0.45 μm capsule filters only					
				, ,		Quick Connector	J	**Single step ½" hose barb and 3/8" Hose Barb end connections are available in 1"capsule filter					ns are no
_					Single Step ½" Hose Barb**	Q	***Male luer slip end connection is available as outlet only in 1" capsule						
						Female Luer Lock	U	filters	sub cua cou		us outlet	01119 111	, capsu
					Male Luer Slip***	W	****3/16" hose barb end connection is available in: - 1" and 2" capsule filters as inlet and outlet - 5" as outlet only						
EFA: Effective Filtration Area			3/16" Hose Barb****	N									
ixample				3/8" Hose Barb**	I	· · · · · · · · · · · · · · · · · · ·							
DWSX		53		01		EE		Х	Х	1		(01

DST DWSX01X1630C

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