

# 0.2 μm *AseptiCap WS-γ* Hydrophilic PVDF Membrane Inline Capsule Filters

**mdi** AseptiCap WS-γ are low protein binding hydrophilic PVDF gamma sterilizable membrane inline capsule filters, designed for sterile filtration of very small fluid volumes in formulation and process development labs.

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.

### **Applications**

#### **Sterile Filtration of**

- Antibodies
- Protein Solutions
- Buffers
- Heat labile additives
- 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

### **Key features**

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

## **Specifications**

		Construction									
Pore Size		0.2μm									
Membrane		Hydrophilic PVDF									
Plastic Compo	nents	Polypropylene									
		Size									
Size		25 mm	50 mm								
Effective Filtrat	ion Area (Nominal)	5 cm <sup>2</sup>	20 cm <sup>2</sup>								
		Integrity Testing/Retention									
Bubble Point (	with Water)	≥ 50psi (3.52 Kg/cm²)									
Microbial Retention		LRV>7 for <i>Brevundimonas diminut</i> a (ATCC 19146) per cm <sup>2</sup>									
		Operational									
Max. Operating Temperature		55 ℃	60 °C								
Max. Differential Pressure		75 psi (5 Kg/cm² @25°C)	42 psi (3 Kg/cm²) @ 30 °C								
Sterilization	By Irradiation	Gamma Irradiatiable up to 50 kGy									
	By Autoclave	Autoclavable at 125°C for 30 minutes, 1 cycle after gamma irradiation. Cannot be in-line steam sterilized.									
		Assurance									
Bacterial Endotoxin		Aqueous extracts exhibit < 0.25 EU/ml as established by Limulus Amebocyte Lysate (LAL) Test as per USP <85>									
Toxicity		Passes Biological reactivity Test, <i>In Vivo</i> , as per USP <88> for Class VI plastics									
Cytotoxicity		Passes Biological Reactivity Tests, In Vitro, USP <87> for cytotoxicity									

Assurance									
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 flushing with a specified volume of WFI								
Extractables with WFI	Passes test as per USP <661>								
Oxidizable Substances	Passes test as per USP <1231>								

### **Ordering Information**

### 25 mm Inline Capsule Filters

Туре		Size		Pore Size		Inlet		Outlet		Radiation Sterilizable		х	Sterility		Pack Size	
	Code		Code		Code		Code		Code		Code			Code		Code
AseptiCap WS	IWSX	25mm	06	0.2 μm	01	1/8" Hose Barb	Н	1/8" Hose Barb	Н	Yes	R		Non Sterile	1	100	04
(0.45µm Upstream)						1⁄4" Hose Barb	В	1/4" Hose Barb	В	No	Х		Gamma Sterile	3		
						Female Luer Lock	М	Male Luer Slip	N							
								Male Luer Lock	L							
Example:																
IWSX		06	5	0	I	M		N		Х		Х	1			04

Note: Gamma Sterile filters can not be sterilized again

### **50 mm Inline Capsule Filters**

Example:

Туре		Size		Pore Size		Inlet		Outlet		Radiation Sterilizable		х	Sterility		Pack	c Size
	Code		Code		Code		Code		Code		Code			Code		Code
AseptiCap WS	. IWSX	50 mm	10	0.2 μm	01	1⁄4" SHB	В	1/4" SHB	В	Yes	R		Non Sterile	1	10	02
(0.45 μm Upstrea	m)					¾" Sanitary Flange	S	¾" Sanitary Flange*	S	No	Х		Gamma Sterile	3	100	04
Vented AseptiCap																

\* In vented AseptiCap WS ¾" Sanitary Flange is available as outlet only

IWSX 10 01