

Surface (Non-woven)



## **MICRO-PURE**

# **AMPX** Type

# Major Applications

Pharmaceutical liquid clarification

Pharmaceutical chemicals, Pharmaceutical ingredients, Drug substance, Manufacturing water

# **Quality standards**

- Manufactured in ISO 9001 certified plant
- FDA 21 CFR compliant
- USP Class VI plastic biological safety testing compliant
- Certificate of quality is attached to the product
- Traceability by lot number

#### **Features**

- 100% Polypropylene
- Pleated type with wide filtration area

# **Advantages**

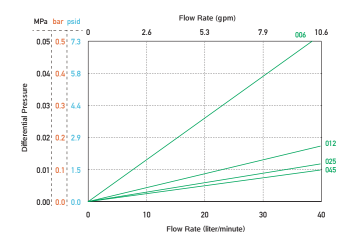
- Wide range of fluid compatibility
- Excellent differential pressure vs. flow rate

#### Specifications

		<u> </u>				
Grades		006	012	025	045	
Micron Ratings ( $\mu$ m)		0.6	1.2	2.5	4.5	
E.F.A. (m²/250L)		0.55	0.75	0.75	0.75	
Media		Polypropylene				
Materials	Core/Cage/Support	Polypropylene				
End Cap		Polypropylene				
Maximum ΔP		0.49MPa at 20℃ (71psi at 68°F)				
Maximum Operating Temp		80℃ (176°F)				
	Length	125/250/500/750 mm				
Dimen- sions	0.D.	70.0mm				
	I.D.	26.1 (for F) / 26.9 (for 3, 4) / 30.0 (for 7) mm				
Available sterilization methods		Inline steam, Autoclave *Applicable to only code 3, 4, 7 with Silicone 0-rings.				
Inline steam sterilization		135°C (275°F) x 30 minutes x 10 cycles *Applicable to only code 3, 4, 7 with Silicone 0-rings.				

## **Differential Pressure vs Flow Rate**

Fluid: Refined Water 20°C (68°F) / Cartridge Length: 250mm



## **Particle Removal Efficiency**

Grades	Particle Removal Efficiency (%)				
Particle Size (μm)	006	012	025	045	
0.6	> 98				
1.2	> 99.9	> 98	> 95	> 95	
2.5		> 99.9	> 99.9	>99.9	

#### Test Conditions

Equipment : Liquid Particle Counter

Filtration : Single Pass Fluid : Refined Water Flow Rate : 10 liter/minute Dust : ACFTD + LATEX Beads

#### Validation items

Items	Evaluation criteria		
Endotoxin (LAL)	Extraction volume with water is less than 0.25 EU/mL and complies with USP 〈85〉 requirements.		
Evaporation residues	Less than 5 mg of evaporation residue per 250 mm cartridge after 24 hours in ultrapure water following autoclave sterilization		
Potassium permanganate consumption	Meets the requirements of the USP Oxidizable Substance Test by flushing with at least 5,000 mL of ultrapure water after autoclaving		
Filter/component toxicity	USP (88) Biological Reactivity Tests For Class VI Plastics compliant		

\*Please refer to the Validation Guide for detailed testing information.

ordering information							
Length	Product Type	Micron Rating	Gasket/0-Ring	End Cap Code	Packaging Code		
5 0 0 L	-AMPX-	0 1 2	S	7	В		
▼		▼	▼	▼	▼		
125 = 125mm		$006 = 0.6 \mu\mathrm{m}$	S = Silicone	F = Flat Gaskets*	A = 1pc		
250 = 250mm		$012 = 1.2 \mu$ m	X = EPDM	3 = 2-222  O-Ring + Fin	B = 6pcs		
500 = 500mm		$025 = 2.5 \mu\mathrm{m}$	T = FEP Encapsulated FKM	4 = 2-222 O-Ring	C = 10pcs		
750 = 750mm		$045 = 4.5 \mu \mathrm{m}$	B = FEP Encapsulated Silicone	7 = 2-226  O-Ring + Fin	F = 25pcs		
				*Code F is only for Silicone			

## **End Cap Code**

Code F

Code 3

Code 4

Code 7

















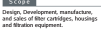
<sup>\*</sup>The performance data listed in the catalog are Typical values obtained under specific conditions based on our tests.





Manufacturing is based on our Quality Management Systems that meet ISO9001 standards.









<sup>\*</sup>The contents of the catalog are subject to change without notice.