

Microcap™ PNYPC

Pharmaceutical Grade
Positively Charged Nylon 6.6
Pleated Membrane Capsules



Microcap™ PNYPC capsule filters consist of a positively charged Nylon 6.6 membrane used for filtering aqueous and non-aqueous liquids that contain negatively charged contaminants.

Available in 0.10, 0.22, 0.45 and 0.65 µm, Microcap™ PNYPC filters are validated for bacteria retention to provide reliable sterile filtration performance.

The positive charge removes negatively charged biological contaminants such as endotoxin, virus and other cell fragments.

Depending on level of contaminant and flow rate, Microcap™ PNYPC filters will typically exhibit > 3-log removal of endotoxin. This combination of functionality makes the PNYPC filter an excellent choice for pharmaceutical and bioprocessing applications.

Typical Applications

Microcap™ PNYPC filters are recommended for sterilising and endotoxin removal in:

- Process water
- Water for injection (WFI)

Features and Benefits

- Validated for use in pharmaceutical applications
- Integrity testable
- Designed for minimal extractables
- Non-fibre releasing
- Low TOC levels
- USP Class VI approved
- Uses FDA complaint materials

Ordering Information

Product Code: 7018 -10-

xxx

x

xx

x

x

Micron Rating (µm)		Pre-sterilised		Length (in)		Inlet		Outlet	
P10	0.10	N	Non-sterile	02	2	A	1/4" Female NPT	A	1/4" Female NPT
P22	0.22	S	Sterile	05	5	B	1/4" Male NPT	B	1/4" Male NPT
P45	0.45			10	10	C	3/8" Female NPT	C	3/8" Female NPT
P65	0.65			20	20	D	1/2" Female NPT	D	1/2" Female NPT
				30	30	E	1/2" Male NPT	E	1/2" Male NPT
						F	1" - 1 1/2" Sanitary	F	1" - 1 1/2" Sanitary
						G	Hose Barb	G	Hose Barb

Specifications

Materials of Manufacture

Housing:	Polypropylene
Filtration media:	Positively Charged Nylon 6,6 Membrane with Polyester support
Media support:	Polypropylene
End caps:	Polypropylene
Centre core:	Polypropylene
Outer support cage:	Polypropylene
Sealing method:	Thermal bonding

Validation

Microcap™ PNYPC filters are validated using test procedures that comply with ASTM F 838-15(ae1) protocols for the determination of bacterial retention in liquids. The challenge level is a minimum of 10^7 organisms per cm^2 of filter media. These capsules have > 7-log removal when challenged with the organisms listed below (0.10 μm and 0.22 μm meet the FDA definition of sterilising grade filters).

0.10 μm :	<i>Brevundimonas diminuta</i>
0.22 μm :	<i>Brevundimonas diminuta</i>
0.45 μm :	<i>Serratia marcescens</i>
0.65 μm :	<i>Saccharomyces cerevisiae</i>

Maximum Operating Parameters

Liquid Operational Pressure	5.52 bar at 20°C (80 psi at 68°F)
Gases Operational Pressure	4.14 bar at 20°C (60 psi at 68°F)
Operating Temperature (water)	43°C at 2.07 bar (110°F at 30 psi)
Reverse Differential Pressure	3.45 bar at 20°C (50 psi at 68°F)
Recommended Changeout Pressure	2.41 bar (35 psi)

Sanitisation and Sterilisation

Autoclave*	121°C (250°F), 30 min, 25+ cycles
Chemical Sanitization	Performed using industry standard concentrations of hydrogen peroxide, peracetic acid and other selected chemicals.

* Note: PNYPC capsules are not designed for steam-in-place (SIP).

Filtration Area

Media	2"	5"	10"	20"	30"
Positively charged Nylon 6,6 Membrane	1.2 ft ² 0.11m ²	3.3 ft ² 0.31m ²	7.0 ft ² 0.65m ²	14.0 ft ² 1.30m ²	21.0 ft ² 1.95m ²

Integrity Testing

Pore Size	Diffusive Flow - Test Pressure *		Minimum Bubble Point *	
	Psi	Bar	Psi	Bar
0.10	48	3.30	**	**
0.22	35	2.41	50	3.5
0.45	20	1.37	25	1.7
0.65	15	1.03	19	1.3

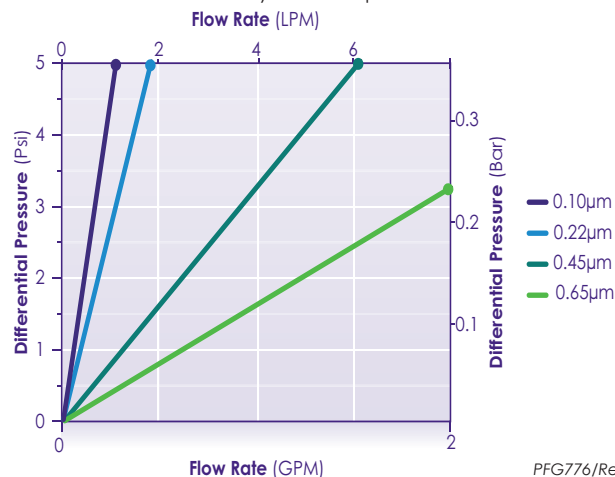
Diffusive Flow Specifications					
Length	2"	5"	10"	20"	30"
m/Lmin	≤ 2.1	≤ 6.3	≤ 15	≤ 30	≤ 45

* For water wetted membrane

** Test pressure exceeds operational limits of capsule filters. Use the Diffusive Flow Test method.

Clean Water Flow Rate

A 2" capsule with 1" sanitary inlet and outlet point, exhibits the flow- ΔP characteristics indicate below, for solutions with a viscosity of 1 centipoise.



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