

Certificate of Quality

The Polyethersulfone Membrane Cartridge Filters have been manufactured in a **mdi** facility in compliance with **ISO 9001** regulations using **validated production processes**.

AseptiSure KS PES Membrane Cartridge Filters

Catalog No. : CPKX5401A0SS101

Type : CPPKS

Pore Size : 0.2 μm (0.45 μm + 0.2 μm)
Lot Number : CK8560C SI.No. 007

SPECIFICATION

| Length | 10" |
|-------------------------------|---|
| Filter Media | Polyethersulfone Membrane |
| Drainage Layers | Polyester |
| Plastic Components | Polypropylene |
| Differential Pressure | < 3.5 Kg/cm² at 25°C |
| Maximum operating Temperature | 80 °C at < 2 Kg/cm² |
| Reverse Pressure | < 0.7 Kg/cm² at 25°C |
| Sterilization | 25 Autoclave/ Steam sterilization cycles at 121°C for 30 minutes each |

LOT RELEASE CRITERIA

100% Integrity Tested : The cartridge filter has been tested for integrity by Air Diffusion Flow test and Bubble

Point test using DI water.

Diffusion flows with DI water were: \leq 30 ml/min @ 2.60 kg/cm² Bubble point value with DI water was: \geq 50 psi (3.44 Bar)

The cartridge filter is also certified for integrity by Bubble point test using 50%

IPA/Water solution.

Bubble point with 50% IPA/Water solution is ≥ 16 psi (1.10 Bar)

Water Flow Rate : ≥ 30 lpm @ 0.70 Kg/cm² @ 27 °C

Microbial Challenge Test : Retains ≥ 10⁷ organisms/cm² of *B. diminuta* ATCC 19146 challenge as per ASTM

F838-05 methodology.

VALIDATED FOR

Heat Stability: Maintains integrity after 25 autoclave/ Steam sterilization cycles at 121°C for 30

minutes each.

Bacterial Endotoxin : Aqueous extracts exhibit < 0.25 EU/mL as established by Limulus Amebocyte

Lysate (LAL) test as per USP <85>.

Extractable: Within limits as specified in USP.

Oxidizable matter: Passes test as per USP.

Biosafety : Passes Biological Reactivity Tests, In Vivo for Class VI plastic as described in USP

<88>.

Cytotoxicity: Passes Biological Reactivity Tests, *In Vitro* as described in USP <87>.

Indirect Food Additives: Passes as per FDA 21CFR 177.1520(a)1(i).

Particle Release : Passes test as per USP <788>, "Particulate matter in Injections".

Fiber Release : Complies with FDA 21CFR 210.3(b)(6).

CUSTOMER SUPPORT

mdi offers its unique interdisciplinary skills to provide solutions to specific problems. Please contact our factory or the local application specialist.

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Head of Quality Assurance Issue Date: 05-Mar-2020 Advanced Microdevices Pvt. Ltd.

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