

STERYKLEAR KSP-KST

- Integrity Testable in situ
- High effective filtering area with SE-TECH technology
- Repeatedly steamable in situ and in autoclave
- Sanitizable
- Thermowelded construction
- EC-listed materials for Food contact
- FDA-listed materials per CFR21
- Bio-Safety per USP-Plastics



STERYKLEAR KSP and KST filter elements adopt SE-TECH tecnology which allows to achieve better filtration results from the membranes; the design optimises the flow distribution between the filter media and the internal core to avoid restrictions and to exploit the full area of the cartridge to generate higher throughput and increase service life.

STERYKLEAR KSP and KST are utilized as final sterilizing filters in pharmaceutical and food & beverage general application; PH-grade, prefluxed with non-pyrogenic water and with certification of quality (with serial number) is used in critical applications. Manufacturing is completed in a controlled environment; secure testing during the manufacturing process assures standards of high quality.

MATERIALS OF CONSTRUCTION

Filter media	Asymmetric PES membrane	
Support KSP	polypropylene	
Support KST	polyester	
Internal Core	polypropylene	
External Cage polypropylen		
End caps / Adapters	polypropylene	

FOOD-SAFETY

STERYKLEAR KSP-KST filter elements meet European Directives 82/711/ECC, 85/572/ECC, 89/109/ECC, 93/8/ECC, 97/48/EC, 2001/61/EC, 2002/16/EC, 2002/72/EC and 2004/19/EC for food contact use and global migration.

BIO-SAFETY

Filter media and components pass USP Biological Reactivity and Chemical-Physical tests for CLASS VI plastics. Specific for "PH" grade: the filter meets USP "Water for injection" requirements for particle release and the effluent is

Non-Pyrogenic per USP Bacterial Endotoxins (< 0,25 EU/ml). **QUALITY STANDARDS**

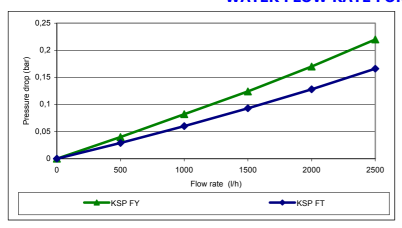
Produced under a certified Quality System to guarantee traceability of manufacturing records and integrity testing results.

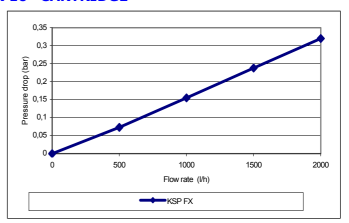
OPERATING CONDITIONS

- max. continuous temperature	80 °C	
- max. cumulative time of steam sterilization	20 hours at 125 °C or 40 hours at 121° C with cycles of 30 minutes	
- sanitization with hot water	80 °C max	
- sanitization with chemicals	can be sanitized by standard chemical agents	
- max. differential pressure	5,0 bar at 25 °C	
- recommended change out differential pressure	e out differential pressure 2,0 bar at 25 °C	
- recommended rinse up volume	3 liters/ 10" cartridge	

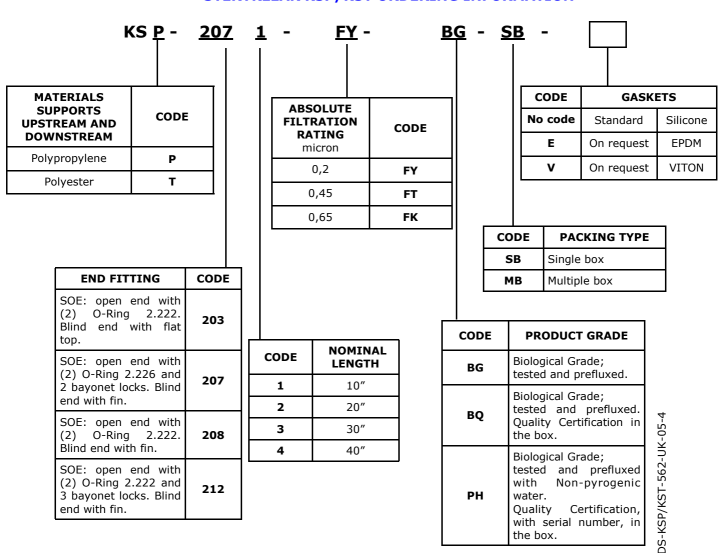
CODE	ABSOLUTE FILTRATION RATING IN LIQUIDS	BACTERIAL RETENTION >10 ¹⁰ CFU/ 10" CARTRIDGE *	ACCEPTABLE LIMIT FOR DIFFUSION TEST WITH WATER FOR 10" CARTRIDGE (ml/min)
FY	0,2 μm	Brevundimonas diminuta	≤ 26 @ 2,7 bar
FT	0,45 μm	Serratia marcescens	≤ 16 @ 1,7 bar
FK	0,65 μm	Leuconostoc oenos	≤ 25 @ 1,1 bar
*as per ASTM F838-83 edition 1993			

WATER FLOW RATE FOR 10" CARTRIDGE





STERYKLEAR KSP/KST ORDERING INFORMATION



Data contained in this bulletin are informative and subject to change without notice. User is responsible for determining whether the product is fit for particular purpose and suitable for User's method of application.



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