

Validation Guide

PVDF Double Layer Hydrophilic Filter Cartridges 0.45+0.2µm

Contents

- 1. Introduction** 1
 - 1.1 Product Description..... 1
 - 1.2 Materials of Construction 1
 - 1.3 Ordering Information..... 2
- 2. Validation Item** 3
- 3. Test Methods and Results** 4
 - 3.1 Correlation of non-destructive integrity testing to liquid bacterial challenge with *Brevundimonas diminuta*(ATCC 19146) for sterilising grade filters 4
 - 3.2 Integrity Test Standard(20°C) 7
 - 3.3 Flow Characteristic(20°C) 7
 - 3.4 Maximum Operating Conditions..... 8
 - 3.5 Steam Sterilization (Thermal Stability)..... 9
 - 3.6 Cleanness..... 10
 - 3.6.1 Gravimetric Extractable..... 10
 - 3.6.2 Fiber Releasing..... 11
 - 3.7 Biological 11
 - 3.7.1 Biological Safety..... 11
 - 3.7.2 Bacterial Endotoxin: LAL Test..... 11
 - 3.8 Chemical Compatibility 12

1.Introduction

1.1 Product Description

Double layer hydrophilic PVDF 0.45+0.2µm filter cartridges are sterilize grade cartridge filters manufactured from asymmetric polyethersulfone membrane and polypropylene hardware providing high flow rates and throughputs, low extractable, broad chemical compatibility. Double layer hydrophilic PVDF 0.45+0.2µm filter cartridges are 100% integrity tested during manufacturing to assure sterilizing-grade performance.

These filter cartridges are designed for removal of particles and the sterile filtration of aqueous solutions.

The double layer hydrophilic PVDF 0.45+0.2µm filter cartridges are designed, developed and manufactured in accordance with an ISO 9001 certified quality management system. These filters are manufactured in a controlled environment that meets the air quality standards of an ISO class 8 room with respect to viable and nonviable particulate and positive pressure.

1.2 Materials of Construction

1.2.1 Component materials used in this product meets the FDA Indirect Food Additive requirements cited in 21 CFR 177-182.

Items	Material
Filter media	hydrophilic PVDF membrane
Support	polypropylene
Cage	polypropylene
Core	polypropylene
End Caps	polypropylene with embedded stainless steel ring
O-ring	silicone











1.2.2 Component materials used in this product meets the regulation (EC) No.1935/2004.

Items	Material
Filter media	hydrophilic PVDF membrane
Support	polypropylene
Cage	polypropylene
Core	polypropylene
End Caps	polypropylene

1.3 Ordering Information

Application	Type	Material	Micron	Length	Adaptor	Sealing	R
D- Dorsan	PVDFPHIL	DL	0.45+0.2µm	5"	E2-222/Flat	S	
Biopharma	Pleated	DL-Double		10"	with SS insert	S-Silicone	
Food&Beverage	Filter Hydrophilic	Layer PES		20"	E3-222/Fin	E-EPDM	
				30"	with SS insert	N-NBR	
				40"	E7-226/Fin	V-Viton	
				with SS insert	F-PTFE		
				E6-226/Flat	F-Encapsuled		
				with SS insert	Viton		
					K-Encapsuled		
					Silicone		

The Catalogue Number is shown as below:

Adaptor No.	Description	Top	Bottom
BLANK SPACE	DOE		
E7	226/Fin with SS insert		
E6	226/Flat with SS insert		
E2	222/Flat with SS insert		
E3	222/Fin with SS insert		

2. Validation Item

Retention	3.1 BCT (ASTM F838-20)	3.2 Integrity Test
Flow Rate	3.3 Flow Characteristic	
Durability	3.4 Maximum Operating Conditions	3.5 Steam Sterilization
Cleanness	3.6.1 Gravimetric Extractables	3.6.2 Non-Fiber Releasing
Biological	3.7.1 Biological Safety	3.7.2 Bacterial Endotoxin
Chemical	3.8 Chemical Compatibility	

3. Test Methods and Results

3.1 Correlation of non-destructive integrity testing to liquid bacterial challenge with *Brevundimonas diminuta* (ATCC 19146) for sterilize grade filters

3.1.1 Introduction

The FDA guidelines on Sterile Products Produced by Aseptic Processing (2004) state, “A sterilizing filter is one which, when challenged with the micro-organism *Brevundimonas diminuta* (*B. diminuta*), at a minimum concentration of 10^7 organisms per cm^2 of filter surface, will produce a sterile effluent”.

In order to meet the requirements of this guideline, liquid challenge tests using *Brevundimonas diminuta* (ATCC 19146) were performed with double layer hydrophilic PVDF $0.45+0.2\mu\text{m}$ filter cartridges using a minimum of 1×10^7 colony forming units (CFU)/ cm^2 of effective filtration area.

The correlation between microbial retention and a non-destructive integrity test is also an important aspect of the validation of sterilizing grade filters. The FDA guideline further states, “After a filtration process is properly validated for a given product, process and filter, it is important to assure that identical filter replacements (membrane or cartridge) used in production runs will perform in the same manner. One way of achieving this is to correlate filter performance data with filter integrity testing data”. The integrity tests used during this validation study were the Forward Flow and Bubble Point tests.

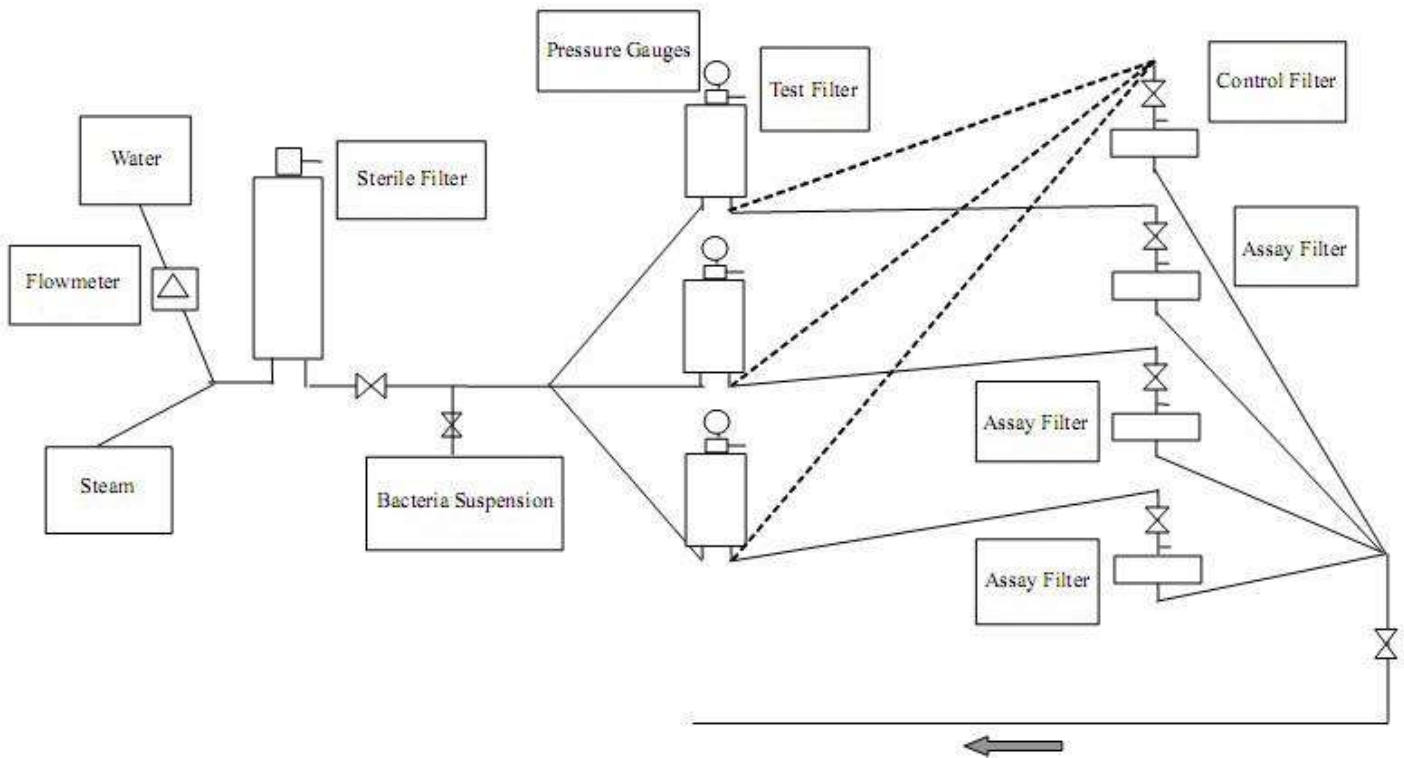
The Forward Flow Integrity Test

In the Forward Flow test, a filter is wetted with a suitable test liquid and a pre-determined gas pressure is applied to the upstream side of the filter assembly. After a stabilization period, the gas flow through the wetted membrane can be measured manually on the downstream side or on the upstream side, using sensitive flow measurement equipment such as the integrity test devices.

The Bubble Point Integrity Test

In the Bubble Point test, a filter is wetted with a suitable test liquid. After a stabilization period, increasing the gas pressure to the upstream side of the filter assembly, using sensitive flow measurement equipment such as the integrity test devices to test the change point of the gas flow rate.

3.1.2 Summary of Methods



Double layer hydrophilic PVDF 0.45+0.2µm filter cartridges with different batch lots were subjected to microbial challenge tests using an aqueous suspension of *Brevundimonas diminuta* (ATCC 19146).

Prior to the challenge tests the filters were installed in an appropriate housing, flushed with DI water at a flow rate of 4 L/min for 15~20 minutes, and then autoclaved at 121 °C for 30 minutes. The filter assembly was then aseptically connected to a pre-sterilized challenge apparatus.

A Forward Flow integrity test was then performed using an integrity test instrument with an air test pressure of 2.7bar (39.1 psi).

An aqueous suspension of *Brevundimonas diminuta* was passed through the filter to achieve a challenge level of $> 1 \times 10^7$ colony forming units (CFU) per cm² of effective filtration area.

During the challenge test, the entire filter effluent was passed through a 0.2 µm-rated analysis disc on the downstream side of the test filter assembly. The filter disc was incubated on TSA and following incubation, the disc was examined to determine if any colonies had grown, indicating whether or not bacteria had passed through the test filter during the challenge.

3.1.3 Test Results

Table listed below indicates double layer hydrophilic PVDF 0.45+0.2µm that are forward flow tested before and after bacterial challenge. The bacterial challenge was conducted using ASTM F838-20 to provide the necessary correlation between a bacterial challenge and a non-destructive integrity test.

Filter type: double layer hydrophilic PVDF 10", 0.45+0.2µm
Challenge organism: *Brevundimonas Diminuta* (ATCC 19146)

Filter Serial Number	Forward Flow mL/min	Challenging bacteria(CFU/10inch)	Sterile Effluent	LRV/cm ²
312180611001	7.2	1.53*10 ¹¹	Yes	7.40
312180611009	7.4	1.95*10 ¹¹	Yes	7.51
312250111002	7.5	2.12*10 ¹¹	Yes	7.54
312250111006	7.6	2.05*10 ¹¹	Yes	7.53
401060511001	7.8	1.53*10 ¹¹	Yes	7.40
401060511007	7.9	1.53*10 ¹¹	Yes	7.40
401060511006	8.1	1.77*10 ¹¹	Yes	7.46
312180611012	8.2	1.95*10 ¹¹	Yes	7.51
312250111005	8.2	1.77*10 ¹¹	Yes	7.46
312180611007	8.3	2.31*10 ¹¹	Yes	7.58
312250111010	8.4	1.95*10 ¹¹	Yes	7.51
312250111012	9.3	2.12*10 ¹¹	Yes	7.54
401060511011	9.5	1.77*10 ¹¹	Yes	7.46
401060511009	9.6	2.31*10 ¹¹	Yes	7.58
401060511010	9.9	2.12*10 ¹¹	Yes	7.54
312250111020	10.2	1.65*10 ¹¹	Yes	7.43
312180611005	10.3	1.19*10 ¹¹	Yes	7.29
312180611006	10.5	2.05*10 ¹¹	Yes	7.53
312250111018	10.8	2.31*10 ¹¹	Yes	7.58
312250111021	11.3	2.82*10 ¹¹	Yes	7.67
312180611006	12.6	1.19*10 ¹¹	Yes	7.29
312180611011	12.8	1.65*10 ¹¹	Yes	7.43
401060511021	12.9	2.05*10 ¹¹	Yes	7.53
401060511013	14.5	2.82*10 ¹¹	Yes	7.67
312180611013	16.1	2.11*10 ¹¹	Yes	7.35
312250111003	16.5	1.65*10 ¹¹	Yes	7.43
401060511004	17.2	2.11*10 ¹¹	Yes	7.31
312250111001	18.9	2.82*10 ¹¹	Yes	7.67
312180611002	21.5	2.11*10 ¹¹	No	<7
312250111004	25.6	1.19*10 ¹¹	No	<7

3.1.4 Conclusions

A Typical double layer hydrophilic PVDF Series 0.45+0.2µm filters from production,

28pcs were found to pass the forward flow integrity test. The table also indicates that double layer hydrophilic PVDF 0.45+0.2µm filter with Forward Flow ≤18.9 mL/min/10" @ 2.7bar at 20°C has sterile filtration efficiency when challenged with > 1 x 10⁷ CFU per cm² of filtration area using *Brevundimonas diminuta*.

Analysis in base of results of forward flow, tested filter ≤18mL/min gave sterile effluent when challenged with LRV>7 of *B.diminuta*.

3.2 Integrity Test Standard(20±5°C)

In base of test results of validation test, forward flow is approved as suitable test methods for double layer hydrophilic PVDF 0.45+0.2µm filter cartridges. We also test the bubble point before and after bacterial challenge. The double layer hydrophilic PVDF 0.45+0.2µm filter with bubble point ≥3.4bar at 20°C has sterile filtration efficiency.

The Integrity test value standard is as below:

Wetting Liquid	DI Water	
Temperature	20±5°C	
Test Gas	Air	
Allowable Integrity Value limit	Forward Flow:	≤18mL/min@2.7bar
	Bubble Point:	≥3.4bar

3.3 Flow Characteristic(20±5°C)

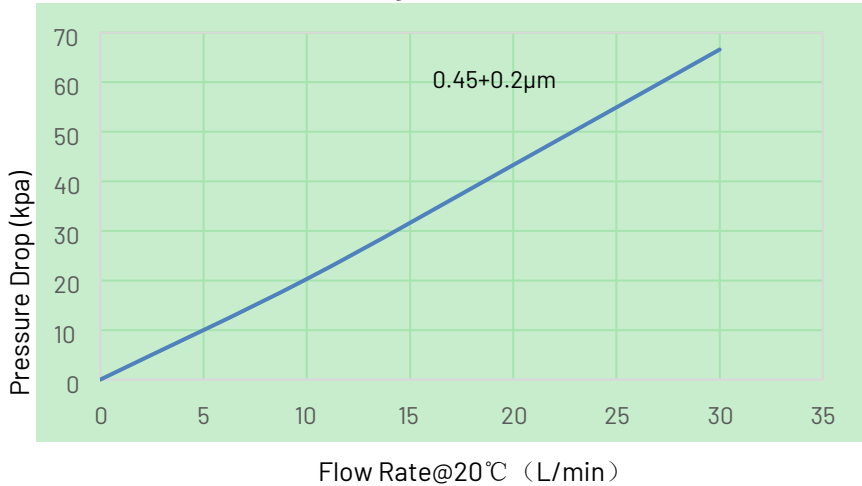
3.3.1 Summary of Methods

Typical double layer hydrophilic PVDF 0.45+0.2µm filter Cartridge from production were used for the tests. The filters were flushed with DI water@2.0bar for15-20mins. Then passed forward flow in base of 2.7bar air test pressure.

Standard production double layer hydrophilic PVDF 0.45+0.2µm filter cartridge (10", EFA 0.6m²) were installed with flow rate test system device, which adjust the flow rate and pressure drop by auto-valve. Then take the continuous records with flow rate, upstream & downstream pressure value, temperature etc.

3.3.2 Test Results

Here is the flow chart of filter cartridge



3.3.3 Conclusions

These data can be used to assist users in sizing filter systems.

3.4 Maximum Operating Conditions

3.4.1 Summary of Methods

Typical double layer hydrophilic PVDF 0.45+0.2µm filter Cartridge from production were used for the tests. The filters were flushed with DI water@2.0bar for 15~20mins. Then passed forward flow in base of 2.7bar air test pressure.

Standard production double layer hydrophilic PVDF 0.45+0.2µm filter cartridge (10", EFA 0.6m²) were installed with pressure drop test system device, which adjust the upstream and downstream pressure by auto-valve.

Use ISO Standard powers to increase the pressure drop till 5.0bar ,temperature 25°C and keep the pressure for 30mins, total 4 cycles.

Use Standard test dust to increase the pressure drop till 2.0bar,temperature 80°C and keep the pressure for 30mins, total 3 cycles. Then test the integrity.

3.4.2 Test Results

Filter Serial Number	Integrity Test (Before the pressure test)		Integrity Test (After the pressure test)	
	Forward Flow(mL/min@2.7bar)	Bubble Point(bar)	Forward Flow(mL/min@2.7bar)	Bubble Point(bar)
401060511002	10.8	4.156	11.1	4.192
401060511005	12.8	3.982	12.9	4.041
312250111007	16.1	3.889	16.3	3.802
312250111011	9.8	4.198	9.9	4.105
312180611004	10.6	4.047	10.7	3.998

The Integrity test value standard is forward flow ≤18mL/min@2.7bar, bubble point ≥3.4bar, test temperature 20±5°C.

3.4.3 Conclusions

Here is the Max. pressure drop and temperature of filter cartridge

Maximum Differential Pressure (Forward)	Maximum Operating Temperature
5.0bar @ 25 °C	80°C
2.0bar @ 80 °C	

3.5 Steam Sterilization(Thermal Stability)

3.5.1 Summary of Methods

The purpose of these tests was to determine the effects of repeated exposure to in-line steam or autoclave cycles on filter integrity using standard double layer hydrophilic PVDF 0.45+0.2µm filters from production. The filters were flushed with DI water @2.0bar for 15~20mins.

Then passed forward flow in base of 2.7bar air test pressure.

During this autoclave study, filters were steamed using high initial differential pressures (1.0bar (14.5 psi)) at 125°C (257°F). The tests were performed in 30 minutes cycles in the forward (out to in) direction.

During the on-line steam sterilization, These filter cartridges were installed in stainless steel housing and steamed in place in the forward(out to in) direction using saturated steam at constant pressure and flow while ensuring effective condensate drainage. After each steam-in-place cycle the filters were cooled by passing dry compressed air through them.

Integrity tests maintained integrity after Sterilized in place and Autoclave to determine the ability of the filter to provide a sterile filtration.

3.5.2 Test Results

Forward Flow (mL/min) after the following Number of 30 minutes Steam Cycles.

Serial No.	0Cycles	10cycles	30cycles	50cycles	100cycles	150cycles
401060511011	11.5	11.7	11.3	11.4	11.2	11.9
401060511012	12.9	12.8	12.5	13.0	12.7	12.8
401060511015	9.7	9.5	9.9	9.6	9.8	9.3
312250111015	8.8	9.1	8.9	8.9	9.1	9.2
312250111016	10.9	11.1	10.8	10.9	11.2	11.1
312250111014	11.8	11.7	12.1	12.4	11.9	12.1
312250111019	12.5	12.8	12.4	12.2	12.1	12.8
312180611010	9.6	9.8	9.9	10.5	9.9	10.1
312180611014	15.4	15.2	15.1	14.8	14.1	14.9
312180611015	12.5	12.9	12.7	13.1	12.8	12.4

The Integrity test value standard is forward flow ≤ 18mL/min @ 2.7bar, test temperature 20 ± 5°C.

Forward Flow (mL/min) after the following number of 30 minutes on-line steam cycles

Serial No.	0Cycles	10cycles	20cycles	30cycles	40cycles	50cycles
312180611016	15.8	16.3	16.5	16.2	16.5	16.2
312180611017	16.6	16.7	16.5	16.9	16.5	16.9
312250111022	14.2	14.7	14.1	14.9	14.1	14.9
401060511016	17.1	17.0	17.5	17.6	17.5	17.6
401060511018	18.2	17.9	18.3	18.6	18.3	18.6

The Integrity test value standard is forward flow ≤ 18mL/min @ 2.7bar, test temperature 20 ± 5°C.

3.5.3 Conclusions

Double layer hydrophilic PVDF 0.45+0.2µm filters have been demonstrated to be capable of withstanding multiple in-line steam/autoclave sterilization cycles.

The data presented in this section support the following product claims for in-line steaming/autoclaving double layer hydrophilic PVDF 0.45+0.2µm filter cartridges:

Sterilized in Place				Autoclave		
Temp.	Time	ΔP	Cycles	Temp.	Time	Cycles
135°C	30min	30kPa	50	125°C	30min	150

3.6 Cleanness

3.6.1 Gravimetric Extractable

3.6.1.1 Summary of Methods

Typical double layer hydrophilic PVDF 0.45+0.2µm filter Cartridge from production were used for the tests.

Preparation of Filter Samples

Extractables tests were performed on typical production filter cartridges (10inch, 0.6m²), which had been autoclaved in order to maximize the quantity of any extractable material present. The filters were wrapped in aluminium foil and autoclaved for half hour at 121°C, using a slow exhaust cycle. Visible droplets of water remaining on the filter elements were allowed to evaporate at room temperature before the extraction was performed.

Extraction Procedure

Dynamic extraction tests were performed. The test filters were immersed in 1800 ml of extraction fluid in a clean measuring cylinder for 24 hours. For four hours the filter was gently moved up and down. This movement created flow through the filter membrane as a result of the pressure head that was created each time the element was partially lifted out of the liquid.

Analysis of Material Extracted

After the extraction, 1500mL of the extraction liquid was evaporated to dryness and the non-volatile extractable were determined gravimetrically.

3.6.1.2 Test Results

Extraction Fluid	Filter serial number	Gravimetric Extractable (milligrams per Filter)	Average(milligrams)
DI Water	312250111024	12.9	15.3
	312250111025	15.4	
	401060511020	17.6	

The Integrity test value standard is forward flow ≤18mL/min@2.7bar, test temperature 20±5°C.

3.6.1.3 Conclusions

The extractable determined of double layer hydrophilic PVDF 0.45+0.2µm filter cartridge were depended by different solvent. The extractable levels under different solvents, different solubility, different temperature and different contact time are not consistent, so it is recommended to test under actual process conditions.

3.6.2 Fiber Releasing

3.6.2.1 Summary of Methods

Typical double layer hydrophilic PVDF 0.45+0.2µm filter cartridges from production were used for the tests.

Filters were autoclaved for one cycle of 30 minutes, 121°C, and then flushed with a total of 10 liters of 0.1 µm filtered water at a flow rate of 1 L per minute. The filtrate was passed through a 0.65µm black gridded disc filter to collect any fibers removed from the filter. Filters were then integrity tested to verify that only integral filters were used in the test.

3.6.2.2 Test Results

double layer hydrophilic PVDF 0.45+0.2µm Filter – Fiber Shedding Results		
Filter serial number	Number of Fibers in filtrate	Forward Flow(mL/min/10" @ 2.7bar, 20 °C)
312180611018	0	14.8
312250111026	0	16.1
401060511022	0	12.0

3.6.2.3 Conclusions

The double layer hydrophilic PVDF 0.45+0.2µm filter cartridges from production don't have Fiber releasing and meet the request of FDA 21 CFR 210.3(b)(6).

3.7 Biological Safety

3.7.1 Biological Tests

3.7.1.1 Summary of Methods

These filters are non-toxic per USP Class VI Biological Tests forPlastics.

Systemic Injection Test, Intracutaneous Test as well as Implantation Test were performed to determine the toxicity of this filters. This testing was performed by an independent laboratory.

3.7.1.2 Conclusions

The materials used in double layer hydrophilic PVDF 0.45+0.2µm filter cartridges from production met the specifications for Biological Reactivity Tests, in vivo, listed in the current revision of the United States Pharmacopeia (USP) for Class VI –121 °C Plastics.

3.7.2 Bacterial Endotoxin: LAL Test

3.7.2.1 Summary of Methods

The test filter was flushed with each of endotoxin-free normal saline .The filtration amount of per filter area is not more than 1mL/cm². Ensure normal saline flow through the entire filter. The aqueous extract was tested with an LAL reagent and all tubes were incubated at 37±1°C for 60±2min.

3.7.2.2 Test Results

Extracts from filters contain <0.25EU/mL endotoxin units per milliliter per the USP Bacterial Endotoxins Test. The results are shown in the following table.

double layer hydrophilic PVDF 0.45+0.2µm filter--Bacterial Endotoxin: LAL Test per USP(+Clotted;-Not Clotted)					
Filter series number	Positive Control	Negative Control	Positive control of test solution	Test solution	Test results (0.25EU/mL)
312180611019	+ +	- -	+ +	- -	<0.25
312180611020			+ +	- -	<0.25
312250111027			+ +	- -	<0.25
312250111029			+ +	- -	<0.25
401060511024			+ +	- -	<0.25
401060511025			+ +	- -	<0.25

3.7.2.3 Conclusions

The double layer hydrophilic PVDF 0.45+0.2µm filter cartridges from production met the specifications for USP Bacterial Endotoxins Test.

3.8 Chemical Compatibility

The chemical compatibility of double layer hydrophilic PVDF 0.45+0.2µm filters is shown in the chart below. Recommendations are based upon static soak for 72 hours at 25°C and 1.0 atmosphere (14.5 psi, 1.01 bar absolute) pressure. Dynamic (operating) conditions at moderate temperatures (±10% fluctuation) will not change the recommendations, but high liquid temperature may do so in some cases.

NOTE: This data is intended to provide expected results when filtration device are exposed to chemicals under static conditions for 48 hours at 25°C, unless otherwise noted, membrane integrity was tested by bubble point.

This chart is intended only as a guide. User should verify chemical compatibility with a specific filter under actual use condition, such as various temperatures, pressure, and concentration.

- R = Resistant.
- L = Limited resistance
- N = Not resistant
- = No data

Chemicals	PVDF Membrane filter	PP Plastic Parts	Silicone O-ring	EPDM O-ring	Viton O-ring
Acetic Acid, glacial	R	R	L	L	N
Acetic Acid, 90%	R	R	L	L	N
Acetic Acid, 30%	R	R	R	L	L
Acetic Acid, 10%	R	R	R	L	L
Hydrochloric acid, conc.35%	R	R	N	N	R
Hydrochloric acid, 20%	R	R	N	N	R
Hydrochloric acid, 3.3%	R	R	-	N	R
Nitric Acid, conc .67%	R	R	N	-	R
Nitric Acid, 27%	R	R	L	L	R
Sulfuric Acid, conc. 96%	R	N	N	-	R
Sulfuric Acid, 16%	R	R	N	-	R
Ammonium Hydroxide 3N,5.7%	R	R	R	-	R
Ammonium Hydroxide 6N,11.4%	R	R	R	-	R
Potassium Hydroxide, 15%	R	R	N	R	R
Sodium Hydroxide 3N, 11%	N	R	R	R	R
Sodium Hydroxide, 22%	N	R	R	R	R
Amyl Alcohol	R	R	N	R	R
Benzyl Alcohol	R	R	L	-	R
Butanol	R	R	L	-	R
Isopropanol	R	R	R	-	R
Methanol	R	R	R	R	N
Ethylene glycol	R	R	R	-	R
Glycerol	R	R	R	R	R
Propylene glycol	R	R	R	-	R
Ethyl ether	R	R	N	N	N
Tetrahydrofuran	N	R	N	N	N
Tetrahydrofuran, 50% v-v	N	R	-	N	N
Acetone	R	R	R	R	N
Cyclohexanone	R	R	L	L	N
Methyl Ethyl Ketone (MEK)	R	R	N	R	N
Methyl Isobutyl Ketone (MIBK)	N	R	N	R	N
Amyl acetate	R	R	N	R	N
Butyl Acetate	R	R	R	-	N
Cellusolve Acetate	R	R	R	-	N
Ethyl Acetate	R	R	L	N	N
Isopropyl acetate	R	R	L	R	N
Methyl acetate	R	R	N	R	N

Carbon Tetrachloride	R	R	N	N	R
Chloroform	R	R	N	N	R
Ethylene dichloride	N	R	N	-	R
Methylene Chloride	R	R	N	N	L
Tetrachloroethylene	R	R	N	-	R
Trichloroethane	N	R	N	N	-
Benzene	R	N	N	N	R
Toluene	R	N	N	N	R
Xylene	R	N	N	N	R
Cottonseed	R	R	R	-	R
Peanut	R	R	R	-	R
Formaldehyde 37%	R	L	L	R	R
Formaldehyde 4%	R	R	R	R	R
Hexan	R	N	N	-	R
Acetonitrile	N	N	N	R	R
Dimethyl Formamide(DMF)	R	R	R	N	L
Dimethylsulfoxide(DMSO)	R	R	N	N	N
Kerosene	R	-	N	N	R
Pyridine	R	L	N	N	N
Petroleum spirits	R	N	N	N	-
Hydrogen Peroxide	N	R	R	R	-
Ozone	N	R	N	L	-
Phenol	-	R	-	-	-

-----END-----