

FilterNYL 0.2 micron

VALIDATION GUIDE

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➤General information about filter cartridges FilterNYL

In a pharmaceutical production process, filtration is a very important step. It guarantees the depletion of any particles or microbial contaminants from the drug being processed. The filtration process must meet high and well defined quality standards.

Filter cartridges validation is mandatory to ensure purity, security and effectiveness to a pharmaceutical product.

The purpose of this document is to supply with useful information every End User of FilterNYL filter cartridges.0.2 µm rated, operating in pharmaceutical applications. The document provides all references about test methods used and results obtained confirming cartridges conformity to the applicable requirements.

FilterNYL filter cartridges 0.2 micron rated, are designed and manufactured for use in production of sterile liquids in pharmaceutical industry. Materials used for the assembly of the cartridges, both the Polytetrafluoroethylene(PTFE) membrane and the Polypropylene polymer used for the hardware supports, had been selected for the specific application meeting the current regulations.

All the assembling steps are carried out by means of thermal bonding processes, without adding adhesive material.. *FilterNYL* filter cartridges 0.2 micron rated, are constructed in controlled contamination clean room and tested prior to the shipment by QA & QC IPM labs, following internal procedures, in order to meet the reference values set.

The identification of the *FilterNYL* filter cartridges is made per the following schema:

NYL	D	10	002	S	I	PH
Product identification NYL	Cartridge type	Lenght	Filtration rating	Gaskets/ O-rings	Insert	Version
	D=DOE(double open end) 2= bayonet 2.226/flat 3= 2.222/flat 5= bayonet 2.222/fin 7= bayonet 2.226/fin 8= 2.222/fin J= Junior cartridge	10=10" 20=20" 30=30" 40=40" 02=2.5" 05=5.0"	002= 0.2 µm 004= 0.45 µm 006 = 0.65 µm 008 = 0.8 µm 012 = 1.2 µm	B= buna N E= EPR S= silicone V= Viton® X= other** ** pl.se contact IPM	I= SS insert* 0= no SS insert * use SS insert when you steam sterilize the filter	PH= Pharma version

Should you need more info, please contact IPM technical support tel. +39 039 21 45 244 or visit the site www.ipmfilters.com

► Filter cartridges FilterNYL 0.2 µm-technical characteristics

Materials of construction				
<i>Membrane</i>	Nylon 6,6 (NYL)			
<i>Support layers</i>	Polypropylene (PP)			
<i>Cartridge supports</i>	Polypropylene (PP)			
<i>O-rings</i>	Silicone			
Technical characteristics				
<i>Filtration area</i>	0.78 m ² (10" module)			
<i>Pores size</i>	0.2 µm			
<i>Length</i>	Cartridges are available with the following dimensions: 5" JUNIOR (125 mm), 10" (250 mm), 20" (500 mm), 30" (750 mm)			
Retention characteristics				
<i>Bacteria retention</i>	Retention of 10 ⁷ CFU/cm ² <i>Brevundimonas diminuta</i> (ATCC 19146), as per ASTM F838-05.			
Mechanical characteristics				
<i>Hydraulic Stress Resistance</i>	<i>FilterNYL filter cartridges 0.2 µm</i> rated showed to retain their diffusion and integrity characteristics after the following hydraulic stress tests			
	Water Pressure (bar)	N° of strokes	Water temperature (°C)	Flow direction
	5.0	50	20±5	Forward
	2.0	50	20±5	Reverse
<i>Thermal Stress Resistance</i>	<i>FilterNYL filter cartridges 0.2 µm</i> rated showed to retain their diffusion and integrity characteristics after the following thermal stress tests			
	Temperature Test (°C)	Pressure Test (bar)	Cycle Time ¹ (hours)	N° of cycles (in autoclave)
	121	1.15	1	50
Extractables				
<i>Extractables</i>	<i>FilterNYL 0.2 filter cartridges meet</i> USP<661>.			
<i>Conformity of materials to the rules for food contact</i>	The materials used to produce the FilterNYL 0.2 µm cartridge filters were confirmed to be compliant with the requirements for food contact prescribed in Reg. N° 1935/2004/EEC and meet the FDA Indirect Food Additive requirements cited in 21 CFR 177 – 182.			
Biological reactivity				
<i>Toxicity of materials</i>	All the materials of the <i>FilterNYL 0.2 µm</i> filter cartridges meet the requirements of USP<87> ("Biological Reactivity Test in vitro for Class VI Plastics") and USP<88> ("Biological Reactivity Test in vivo for Class VI Plastics – Safety Test").			
Release of particles and/or Endotoxins				
<i>Bacterial endotoxins</i>	< 0.125 EU/mL of aqueous extract as per USP<85>.			
<i>Non-fibre releasing</i>	FilterNYL 0.2 µm filter cartridges satisfy the requirements as per USP<788>			

¹ Total time considering warming up time of 15 minutes; 30 minutes at test temperature and 15 minutes for cooling down to ambient temperature.

►Bacteria removal efficiency

Regulator Authorities FDA and EMA, set the rule to demonstrate the sterilizing action of a filter before to start to use it in pharmaceutical applications. So, the sterilizing grade of a filter cannot be set only by the rating of pores dimension.

In order to consider a filter of sterilizing grade it must be submitted to a process simulation challenging it by a bacterial suspension of *Brevundimonas diminuta* ATCC 1946 at the concentration of not less than 10^7 CFU/cm². The downstream fluid must be sterile.

The correlation between the microbial retention and the non-destructive integrity test is another very important aspect of the validation of a pharmaceutical grade filter cartridge.

The Regulator Authorities require a procedure to control the filter both prior and after the use.

Hereinafter are reported the Challenge tests results of tests made on FilterNYL filter cartridges 0.2 µm rated. According to the ASTM F838-05 method, the tests demonstrate the ability of membranes, used to produce FilterNYL, to fully retain a *Brevundimonas diminuta* bacterial charge of 10^7 CFU/cm².

You can also find the correlation between the retention and the diffusion test.

This correlation is important to set the maximum diffusion flow acceptance limit to guarantee that the filter will produce a sterile effluent.

In the following tables are reported the correlation between the diffusion test (non destructive test) and the bacterial challenge test (destructive test) for FilterNYL 0.2 µm filter cartridges Part Number: NYL J05 002 SI - PH ; NYL 710 002 SI - PH and NYL 7 20 002 SI - PH.

►Filter Cartridge type NYL J05 002 S I - PH 0.2 micron rated Challenge test

To perform the test, 20 filter cartridges were withdrawn from normal production. Each cartridge has been tested by diffusion test and then challenged by a suspension of *Brevundimonas diminuta* at the concentration $>10^7$ CFU/cm². Numerical and graphical results are reported in Table 1 and Figure 1 respectively. The filter cartridges NYL J05 002 S I – PH meet acceptability criteria on performance basis.

Lot N	Diffusion at 2200 mbar (mL/min)	Downstream Sterility	Titre Reduction
6389/31	12.0	Yes	$> 1.5 \times 10^8$
6161/12	12.5	Yes	$> 1.6 \times 10^8$
5967/02	13.0	Yes	$> 1.4 \times 10^8$
5876/17	13.5	Yes	$> 1.5 \times 10^8$
5885/28	13.8	Yes	$> 1.6 \times 10^8$
6389/28	13.9	Yes	$> 1.7 \times 10^8$
6399/11	14.0	Yes	$> 1.7 \times 10^8$
6161/12	14.0	Yes	$> 1.9 \times 10^8$
6377/18	14.2	Yes	$> 1.4 \times 10^8$
5967/02	14.3	Yes	$> 1.5 \times 10^8$
5876/17	14.3	Yes	$> 1.7 \times 10^8$
6111/10	14.4	Yes	$> 1.8 \times 10^8$
5992/01	14.5	Yes	$> 1.8 \times 10^8$
6161/19	14.5	Yes	$> 1.4 \times 10^8$
5992/32	14.8	Yes	$> 1.7 \times 10^8$
6111/22	14.8	Yes	$> 1.7 \times 10^8$
5992/14	15.0	Yes	$> 1.8 \times 10^8$
5975/20	15.0	Yes	$> 1.2 \times 10^9$
5975/26	15.5	No	1.2×10^6
5834/12	16.0	No	1.8×10^6
5832/07	16.5	Yes	$> 1.8 \times 10^8$

Table 1- FilterNYL: NYL J05 002 S I - PH filter cartridges 0.2 µm rated
Results of bacterial retention test

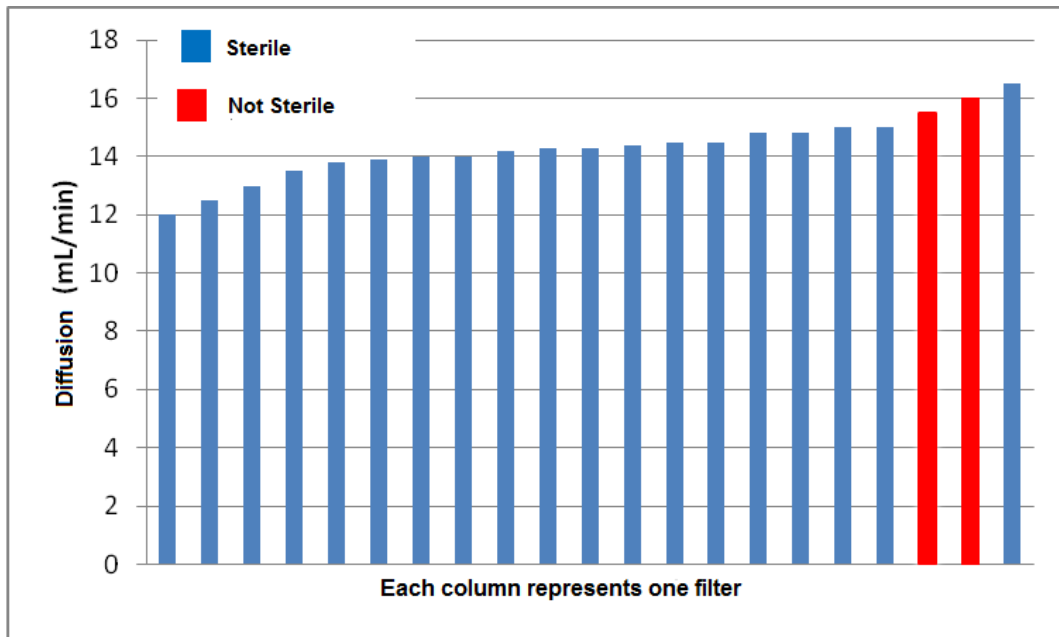


Figure 1- FilterNYL: NYL J05 002 S I - PH filter cartridges 0.2 μ m rated. Correlation between diffusion test and bacterial retention test. ASTM F838-05

On performance basis, FilterNYL filter cartridges NYL J05 002 S I – PH 0.2 micron rated meet acceptability criteria and are considered to be a sterilizing filters within a maximum diffusion value of 15.0 mL/min

►Filter cartridge type NYL 7 10 002 S I – PH: correlation diffusion vs retention

To perform the test, 20 filter cartridges were withdrawn from normal production. Each cartridge has been tested by diffusion test and then challenged by a suspension of *Brevundimonas diminuta* at the concentration $>10^7$ CFU/cm². Numerical and graphical results are reported in Table 2 and Figure 2 respectively. The filter cartridges NYL 7 10 002 S I – PH meet acceptability criteria on performance basis.

LOT N°	DIFFUSION AT 2200 mbar (ml/min)	DOWNSTREAM STERILITY	TITLE REDUCTION
6634/31	25.5	Yes	$> 1.4 \times 10^8$
6634/22	26.0	Yes	$> 1.4 \times 10^8$
6213/12	26.5	Yes	$> 1.2 \times 10^8$
6114/07	27.0	Yes	$> 1.6 \times 10^8$
5974/12	27.5	Yes	$> 1.7 \times 10^8$
5764/30	28.0	Yes	$> 1.8 \times 10^8$
5764/12	28.1	Yes	$> 1.8 \times 10^8$
5949/33	28.4	Yes	$> 1.4 \times 10^8$
5764/18	28.4	Yes	$> 1.5 \times 10^8$
5935/35	28.7	Yes	$> 1.5 \times 10^8$
5882/05	28.9	Yes	$> 1.3 \times 10^8$
6435/26	29.0	Yes	$> 1.4 \times 10^8$
6465/19	29.3	Yes	$> 1.2 \times 10^8$
6354/14	29.3	Yes	$> 1.1 \times 10^8$
6215/37	29.5	Yes	$> 1.4 \times 10^8$
6215/34	29.8	Yes	$> 1.7 \times 10^8$
6215/02	30.0	Yes	$> 1.8 \times 10^8$
6632/46	30.0	Yes	$> 1.3 \times 10^9$
6176/17	30.5	No	1.6×10^6
6989/12	31.0	No	1.5×10^6
6045/22	33.0	Yes	$> 1.7 \times 10^8$

Table 2- Filter NYL: NYL J05 002 S I - PH filter cartridges 0.2 μ m
Results of bacterial retention test

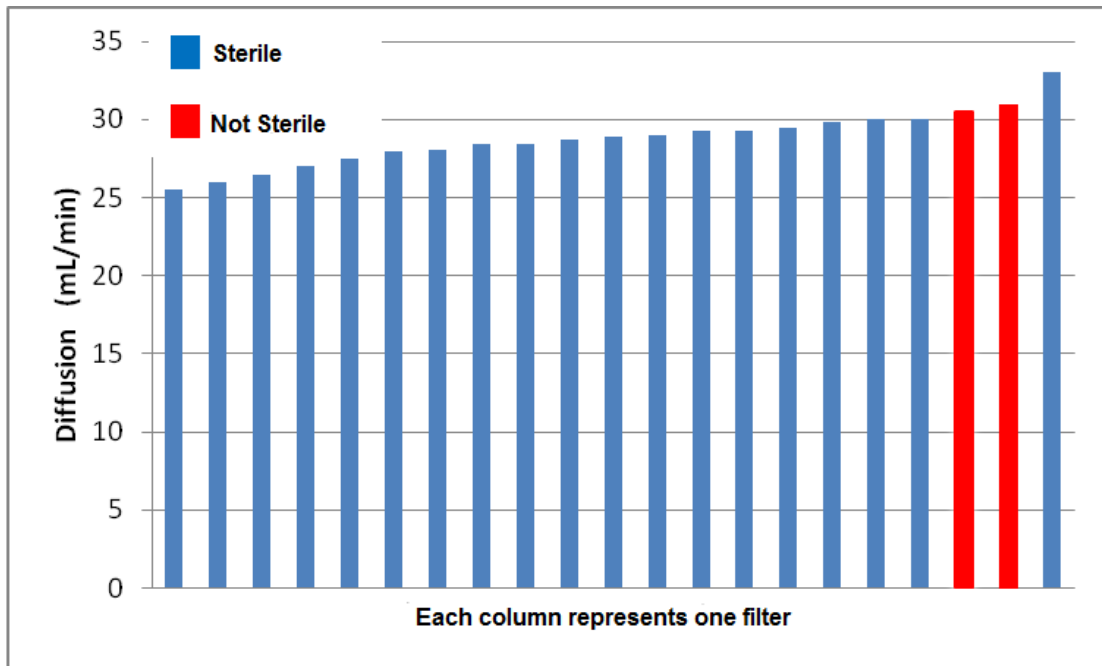


Figure 2- FilterNYL: NYL 7 10 002 S I - PH filter cartridges 0.2 μ m rated. Correlation between diffusion and bacterial retention test. ASTM F838-05

On performance basis, FilterNYL filter cartridges NYL 7 10 002 S I - PH 0.2 micron rated meet acceptability criteria and are considered to be a sterilizing filters within a maximum diffusion value of 30.0 mL/min.

► Filter cartridge type NYL 7 20 002 S I – PH correlation diffusion vs retention

To perform the test, 20 filter cartridges were withdrawn from normal production. Each cartridge has been tested by diffusion test; and then challenged by a suspension of *Brevundimonas diminuta* at the concentration $>10^7$ CFU/cm². Numerical and graphical results are reported in Table 3 and Figure 3 respectively. The filter cartridges NYL 7 20 002 S I – PH meet acceptability criteria on performance basis.

LOT N°	DIFFUSION AT 2200 mbar mL/min	DOWNSTREAM STERILITY	TITLE REDUCTION
4833/23	56.0	Yes	$> 1.5 \times 10^8$
4976/27	56.5	Yes	$> 1.6 \times 10^8$
4976/15	57.0	Yes	$> 1.5 \times 10^8$
4976/33	57.5	Yes	$> 1.8 \times 10^8$
4833/16	57.5	Yes	$> 1.8 \times 10^8$
4987/18	57.7	Yes	$> 1.8 \times 10^8$
4961/39	57.9	Yes	$> 1.7 \times 10^8$
5234/03	58.1	Yes	$> 1.9 \times 10^8$
5234/12	58.4	Yes	$> 1.3 \times 10^8$
5234/09	58.5	Yes	$> 1.3 \times 10^8$
4885/25	58.7	Yes	$> 1.5 \times 10^8$
4885/05	58.8	Yes	$> 1.5 \times 10^8$
5334/32	59.0	Yes	$> 1.9 \times 10^8$
5668/25	59.2	Yes	$> 1.3 \times 10^8$
5100/05	59.6	Yes	$> 1.5 \times 10^8$
5810/31	59.8	Yes	$> 1.6 \times 10^8$
5810/35	60.0	Yes	$> 1.6 \times 10^8$
5973/03	60.0	Yes	$> 1.3 \times 10^9$
5964/15	60.5	No	1.8×10^6
5967/18	61.0	No	1.6×10^4
5234/15	62.5	Yes	$> 1.7 \times 10^8$

Table 3- Filter cartridges FilterNYL: NYL 7 20 002 S I – PH, 0.2 µm rated
Bacterial retention test results

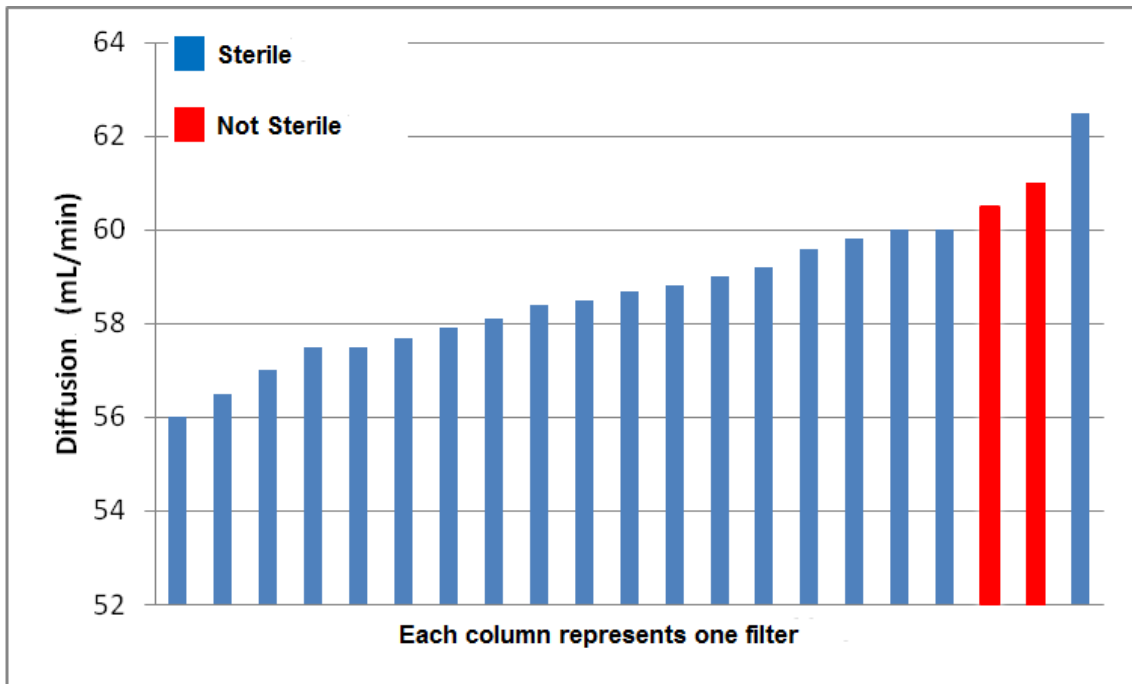


Figure 3- FilterNYL filter cartridges NYL 7 20 002 S I – PH 0.2 micron rated. Correlation between diffusion and microbial retention test as per ASTM F838-05

On performance basis, FilterNYL filter cartridges NYL 7 20 002 S I - PH 0.2 micron rated meet acceptability criteria and are considered to be a sterilizing filters within a maximum diffusion value of 60.0 mL/min.

Biological and extractables tests

►Bacterial endotoxins – current USP <85>

The test was used to measure bacterial endotoxins released from FilterNYL filter cartridge 0.2 micron rated. The test was carried out according to the Current USP<85> using the Chromogenic Method to measure the amount of endotoxins released by a filter cartridge type FilterTEF 0.2 micron.

The sample cartridge was filled with a pyrogen-free water and then treated by an ultrasonic device in order to enhance endotoxins extraction.

Water samples obtained was then submitted to the analysis.

The lab test shows bacterial endotoxins level released by the filter cartridge FilterTEF.

The test results demonstrate the absence of endotoxins release from FilterNYL cartridges.

LOT N°	METHOD	RESULTS
5967/03	cUSP<85>	<0.25 EU/mL
5876/21	cUSP<85>	<0.25 EU/mL
6111/15	cUSP<85>	<0.25 EU/mL

Table 4-LAL test results

►Chemical compatibility

The following table lists chemical compatibility of FilterNYL filter cartridge 0.2 micron rated, when challenged with chemical products. Chemical compatibility is influenced by several parameters (temperature, concentration, contact time). IPM suggests that the Customers check chemical compatibility running a test at the same process conditions, before to use a FilterNYL cartridge 0.2 micron. IPM remains available, if requested, to supply the Customer with technical support..

CHEMICAL CLASSIFICATION	COMPOUND (Test Conditions 7days at 20°C)	RESULTS
ACIDS	Acetic acid, 90%	N
	Acetic acid, 30%	N
	Acetic acid, 10%	N
	Hydrochloric acid, conc. (35%)	N
	Hydrochloric acid, 6N (20%)	N
	Hydrochloric acid, 1N (3.3%)	N
	Nitric Acid, conc. (67%)	N
	Nitric acid, 6N (27%)	N
	Sulphuric acid, conc. (96%)	N
	Sulphuric acid, 6N (16%)	N
ALCOHOLS	Amyl alcohol	R
	Benzyl alcohol	L
	Butyl alcohol	R
	Ethanol (Ethyl alcohol)	R
	Isopropanol	N
	Methanol	R
AROMATIC HYDROCARBONS	Benzene	R
	Toluene	R
	Xylene	R
BASES	Ammonium hydroxide, 3N (5.7%)	R
	Ammonium hydroxide, 6N (11.4%)	R
	Potassium hydroxide, 3N (15%)	R
	Sodium hydroxide, 3N (11%)	R
	Sodium hydroxide, 6N (22%)	R
ESTERS	Amyl acetate	R
	Butyl acetate	R
	Cellosolve acetate	R
	Ethyl acetate	R
	Isopropyl acetate	R
	Methyl acetate	R
ETHERS	Ethyl ether	R
	Tetrahydrofuran (THF)	R
GLYCOLS	THF in water (50:50)	R
	Ethylene glycol	R
	Glycerol	R
HALOGENATED HYDROCARBONS	Propylene glycol	R
	Carbon tetrachloride	R
	Chloroform	R
	Ethylene dichloride	L
	Methylene chloride	L
KETONES	Acetone	L
	Cyclohexanone	R
	Methyl ethyl ketone (MEK)	R
	Methyl isobutyl ketone	R
OILS	Cottonseed	R
	Peanut	R
MISCELLANEOUS	Acetonitrile	L
	Dimethyl-Formamide (DMF)	L
	Formaldehyde (37%)	R
	Hexane	R
	Pyridine	N

Table 5 - FilterNYL 0.2 micron cartridges Chemical Compatibility
(R=recommended; L=Limited resistance; N=Not Recommended; « - » Insufficient Data)

➤Extractables – current USP<661>

Test scope is to measure the amount of non-volatile residual released by the cartridge treated with WFI water. The extraction tests were run in static conditions for 24 hours at 70 °C. A portion of liquid extracted was evaporated to dryness and then submitted to non-volatile residual (NVR) measure.

NVR test results are reported in table 4.

LOT N°	NVR
5967/05	<5 mg
5876/22	<5 mg
6111/16	<5 mg

Table 6-Results of tests carried out according to cUSP<661>

FilterNYL 0.2 micron meet USP<661> requirements.

Biological Safety Test

►"In vitro" Biological Reactivity Test - Class VI Plastic Materials (current USP<87>)

Test scope is to evaluate biological toxicity of FilterNYL cartridges 0.2 micron . The cartridge (O-rings included) is submitted to extraction with extractive solutions following related recipes provided for by current USP <87>. Test results are recorded as reactivity degree ranging from 0 (none reactivity) to 4 (high reactivity)

LOT N°	TEST	METHOD	RESULTS
5876/25	cytotoxicity	cUSP <87>	Reactivity grade = 0
6111/01	cytotoxicity	cUSP <87>	Reactivity grade = 0
5831/53	cytotoxicity	cUSP <87>	Reactivity grade = 0

Table 7-Results of tests carried out according to current USP<87>

Test results carried out according to USP<87>,.demonstrate lack of "in vitro" biological reactivity by FilterNYL filter cartridges 0.2 micron rated.

► **“In Vivo” Biological Reactivity Test - Class VI Plastic Materials – Safety Test (current USP<88>)**

Test scope is to evaluate the “in vivo” reactivity of FilterNYL cartridge according to the method provided for by USP <88>.

Test results are compared with USP acceptability limits.

The results are reported in the table.

LOT N°	TEST	RESULTS
5876/01	Intracutaneous Test	Conform
	Systemic Injection Test	Conform
	Implantation Test	Conform
6111/19	Intracutaneous Test	Conform
	Systemic Injection Test	Conform
	Implantation Test	Conform
5831/11	Intracutaneous Test	Conform
	Systemic Injection Test	Conform
	Implantation Test	Conform

Table 8-Results of the tests carried out according current USP<88>

Test results carried out according to USP<88>, demonstrate lack of “in vvo” biological reactivity by FilterNYL filter cartridges 0.2 micron rated.

Physical characteristics

►Hydraulic stress test

Test purpose is to demonstrate the capability of FilterNYL 0.2 micron cartridges to tolerate hydraulic stress at ambient temperature (20 ± 5 °C). 10" FilterNYL cartridges were installed inside an holder, wetted with water and fluxed at 2 bar during 20 minutes at least. After the wetting cycle was run a diffusion test.

If the cartridge passes the diffusion test, is then washed with a direct flux during 10 minutes, with 50 sudden pressure strokes from 0 to 5.0 bar. The same cartridge is then back-flushed at 2.0 bar per 10 minutes, with 50 sudden pressure strokes from 0 to 5.0 bar.

This test is repeated 50 times in both flux directions- Every 10 stress cycle every single cartridge is controlled by integrity diffusion test.

Diffusion test is performed at 2200 mbar. Acceptability limit of diffusion test was set at 30.0 mL/min. The following table reports diffusion test results run before and after the stress test.

LOT N°	DIFFUSION BEFORE STRESS TEST (mL/min)	DIFFUSION AFTER STRESS TEST (mL/min)
6633/41	28.0	27.5
6213/37	28.5	29.0
6114/19	27.5	29.0
5974/35	28.5	29.0
5764/09	29.0	28.5
5935/07	28.0	28.5
5831/45	28.0	29.5
5994/31	28.0	29.0
6213/50	27.0	29.0

Table 9-Results of hydraulic stress test run on FilterNYL filter cartridges

Test results confirm the high resistances of FilterNYL filter cartridges to hydraulic stress, even changing pressure and flux direction.

► Thermal stress test

The purpose of this test is to determine FilterNYL filter cartridges (10") resistance to multiple steam sterilization. The test has been carried out at 121 °C as provided for an appropriate steam sterilization. After a diffusion test the cartridge is submitted to 50 sterilization cycles. Every 10 cycles diffusion test is repeated.

Sterilization cycles follow the schedule below

STEP	TEMPERATURE (°C)	PRESSURE (bar)
STERILIZATION	121±1	1.14±0.04
WARM UP	121→96	1.12→0.00

Table 10 -Sterilising cycle program

Diffusion tests were run at 2200 mbar. Acceptability limit of diffusion test was set at 30.0 mL/min. The following table reports diffusion test results run before and after the stress test:

LOT N°	DIFFUSION BEFORE STRESS TEST (mL/min)	DIFFUSION AFTER STRESS TEST (mL/min)
6633/44	28.5	29.0
6213/26	29.0	29.5
6114/13	27.5	29.5
5974/11	28.0	29.0
5764/01	28.0	29.5
5935/54	29.0	28.0
5831/32	28.5	28.0
5994/12	29.5	29.5
6213/02	27.5	28.5

Table 11-FilterNYL cartridges Thermal Stress Test results

Test results confirm the high resistances to thermal stress of FilterNYL filter cartridges submitted to repeated thermal stress in autoclave.

► Differential pressure vs water flow rate

The test measures the pressure drop when different flow rates are fluxed across the filter cartridge membrane.

All data are referred to temp of 20±5 °C.

10" FiterNYL cartridges typical pressure drop vs flow rate curve, is shown in the following graph.

Within the typical flow rate range of usage, FilterNYL cartridges guarantee initial pressure drops less than 0.2 bar (flow rate <1200 L/h per 10" cartridge)

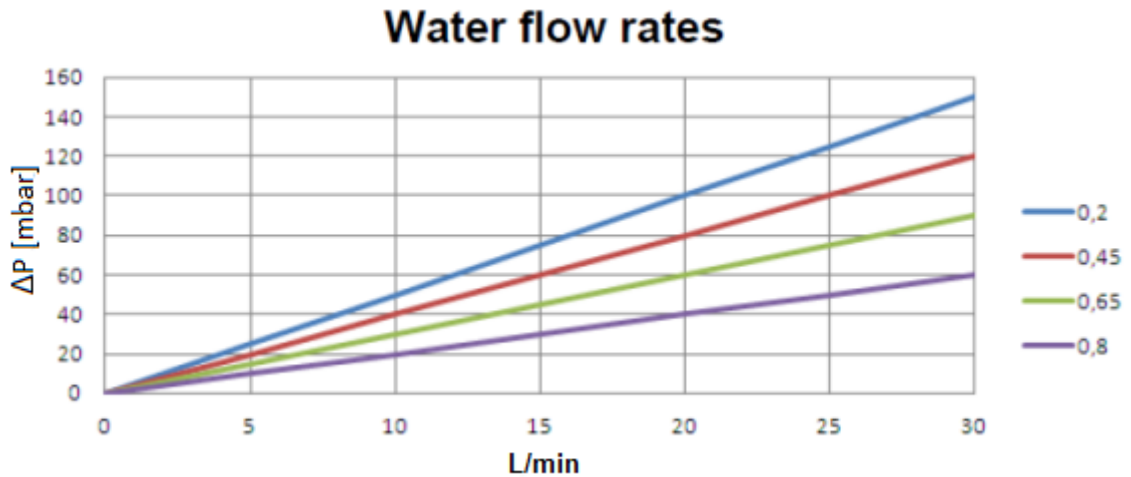


Figure 4-Water Flow Rate vs pressure drop graph.

NYL	D	10	002	S	I	PH
	Cartridge type	Lenght	Filtration rating	Gaskets/O-rings	Insert	Version
Product identification NYL	D=DOE(double open end) 2= bayonet 2.226/flat 3= 2.222/flat 5= bayonet 2.222/fin 7= bayonet 2.226/fin 8= 2.222/fin	10=10" 20=20" 30=30" 40=40"	002= 0.2 μm 004= 0.45 μm 006 = 0.65 μm 008 = 0.8 μm 012 = 1.2 μm	B= buna N E= EPR S= silicone V= Viton® X= other** ** pl.se contact IPM	I= SS insert* 0= no SS insert * use SS insert when you steam sterilize the filter	PH= Pharma version
	J= Junior cartridge	02=2.5" 05=5.0"				

Attachment

► Sample of Certificate of Conformity (CoC) enclose in every packaging of cartridge type –PH

Every cartridge type –PH (which can be used in pharmaceutical productions) is shipped together with a Conformity Certificate (CoC) / Lot traceability. Here following a facsimile:

 <p>Via Melegnano, 22 20035 Sesto San Giovanni - Italy Tel. +39 039 2145244 Fax +39 039 2145245 E-mail: ipm@ipmsistemi.com www.ipmfilters.com</p>		<p><i>Data di produzione</i> <i>Date of production</i> <i>Date de production</i> <i>Fecha de producción</i></p> <p>01/01/2013</p>	
<p>CERTIFICATO DI CONFORMITÀ CERTIFICATE OF CONFORMITY CERTIFICAD DE CONFORMITAT CERTIFICADO DE CONFORMIDAD</p>		<p>XXXXXXXX</p>	
<p><i>Cliente - Customer</i> <i>Cliet - Cliente</i></p>		<p>XXXXXXXXXXXXXXXXXXXX</p>	
<p><i>TIPO DI FILTRO</i> <i>FILTER TYPE</i> <i>TYPE DE FILTRE</i> <i>TIPO FILTRO</i></p>		<p>Filter NYL</p>	
<p><i>MOD.</i></p>		<p>NYL710.002-S1</p>	
<p><i>LOT.</i></p>		<p>L/N: XXXX</p>	
<p><i>Controllo visivo e dimensionale:</i> <i>Visual and dimensional check:</i> <i>Contrôle visuel et dimensionnel:</i> <i>Inspección visual y control dimensional:</i></p>		<p>✓</p>	
<p><i>Risultato - Result - Résultat - Resultado:</i></p>		<p>ACCEPTED</p>	
<p><i>Test di integrità (diffusione):</i> <i>Integrity test (diffusion):</i> <i>Test d'intégrité (diffusion):</i> <i>Prueba de integridad (difusión):</i></p>		<p>✓</p>	
<p><i>Limite di accettabilità:</i> <i>Accepted values:</i> <i>Limite de tolérance:</i> <i>Valor límite aceptable:</i></p>		<p>As per IPM QC procedures</p>	
<p><small>Il test di integrità come sopra specificato, è stato eseguito su ogni singolo cartuccia di questo lotto e risponde alle procedure interne del controllo qualità. IPM certifica che le cartucce filtranti IPM sopra descritte sono state controllate in ambiente controllato. Materia di costruzione: L'impregnato del filtro rispondente alle specifiche per test di biocompatibilità previste dalla USP classe VI per materie plastiche a 121°C su quanto previsto per filtri a contatto con alimenti secondo CFR Title 21 parte 170-199. Questo prodotto risponde a tutti gli standard qualitativi IPM. Il prodotto non è sterile. IPM dichiara che questo certificato di conformità riguarda il reso e sostituisce quello del Vostro cliente. Tutti i test sopra descritti sono stati condotti secondo procedura. La merce ed i materiali sono privi di difetti. This test has been run on every cartridge of this lot and meets the internal QC procedures. IPM hereby certifies that the above IPM filter cartridge has been manufactured in a controlled environment. Material of construction: The filter components have met the specifications for biocompatible tests listed in the current version of the United States Pharmacopoeia (USP) for Class VI plastics at 121°C. These filters also are made from materials listed for head contact usage per 21 CFR 171 of the U.S. Code of Federal Regulation (CFR) parts 170-199. Contact IPM for more information regarding materials of construction. The filters are not supplied sterile. IPM declares that this certificate of conformity is true and correct and meets your order requirements. All the above tests were performed as per our procedures and are to certify grade and materials have no defect. Le test d'intégrité de diffusion est effectué sur chaque cartouche du lot et répond toutes les exigences internes du contrôle de qualité. IPM certifie que les cartouches filtrantes IPM ci-dessus décrites ont été fabriquées en milieu contrôlé. Matière de construction: Les composants du filtre correspondent aux critères des tests de biocompatibilité prévus par l'USP classe VI pour matières plastiques à 121°C définies pour les liquides en contact alimentaire selon le 21 CFR partie 170-199. Ce produit n'est pas stérile. IPM déclare que ce certificat de conformité est vrai et correct et remplit toutes vos exigences. Tous les tests décrits ci-dessus sont conformes à la procédure. La marchandise et les matériaux n'ont aucun défaut. Todos y cada uno de los cartuchos de este lote han sido sometidos al test de integridad según las especificaciones antes descritas, y cumplen con los procedimientos internos del control de calidad. IPM certifica que las cartuchas filtrantes IPM antes descritas han sido fabricadas en ambiente controlado. Materiales de fabricación: Los componentes del filtro cumplen los criterios para pruebas de biocompatibilidad de la USP clase VI para plásticos a 121°C y con todos los requisitos para filtras en contacto con alimentos según el título 21 de CFR partes 170-199. Este producto no es estéril. IPM declara que los datos de este certificado de conformidad son verídicos y se corresponden con los requisitos de su pedido. Todos los pruebas antes citadas se han llevado a cabo con arreglo a sus procedimientos. Los productos y los materiales están libres de defectos.</small></p>			
<p>IL CONTROLLO DI QUALITÀ - QUALITY CONTROL CONTRÔLE DE QUALITÉ - CONTROL DE CALIDAD</p>			

Figure 5- Sample of Conformity Certificate released by IPM Sistemi di Filtrazione

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Doc n° VGNYL02EN130003 rev. 00 printed in Italy 02/2014

IPM reserves the right to change any and all characteristic of the items to offer the best product to the customer. Please check with IPM if any doubt about.

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