Tofflon Life Science



Filtration Product

Tofflon Life Science

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Website

LinkedIn

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Filter Plant Introduction

The filter manufacturing plant of Tofflon Life Science located in Hangzhou, and is one of the core production site in the field of filtration and separation technology of Tofflon Group. The plant is dedicated to providing stable and high-quality filter consumables for biopharmaceutical industry.

The plant has 12,000 m² in space and has a validation center which has passed CNAS certificate (ISO17025). The plant can not only provide filters to customer but also provide related validation service at the same time. It has four kinds of major filter consumables include virus filter, TFF cassette, depth filer and sterile filter which cover all downstream filtration process in biopharma production, makes its capability to provide one-stop solutions for downstream protein separation and purification.

The existing products include:

Valpha® virus removal filter, Vshell® virus removal prefilter, TanFlux® II TFF cassette, Saiclear® depth filter, Saipress® sterilization-grade hydrophilic filter, Saipress® HF sterilization-grade hydrophilic filter, Saipress® PVDF hydrophilic sterile filter, Filgiant® sterilization-grade hydrophobic filter, Filgiant® L series hydrophobic filter, Felix 6KP integrity tester, Saiflow™ pharmaceutical-grade stainless steel housing, PolySai™ polypropylene prefilter, peristaltic pump and torque wrench.

Valpha® Virus Retention Membrane

Biosafety Security for Your Virus Removal Process

Tofflon Valpha® family of virus removal filters are designed to efficiently remove virus particles from biological products (plasma products, antibody drugs, etc.). The Valpha® family of virus removal filters are validated to provide you reliable virus retention even at higher levels of virus challenge.

Key to virus security

Valpha® virus removal filters use membrane filtration technology to remove virus particles based on size exclusion. It uses an innovative high-performance polymer membrane to trap the virus on the membrane surface and in the membrane pores. Key performance parameters are usually logarithmic removal rate (LRV), flux, and load. These parameters are influenced by many interacting factors, such as virus loading, protein concentration, impurity concentration, pressure, process flow rate, ion strength, and process interruption. Therefore, it is very important to select and validate the appropriate virus removal filter under the established process conditions.

Valpha® family virus removal filters provide you two choices of membrane: Valpha RC(Regenerated Cellulose) and Valpha PES (Modified PolyEtherSulfone).

Valpha® RC HP uses regenerated cellulose membrane. Cellulose is a high hydrophilic material and is widely used as a basic polymer in the preparation of filtration membranes. The natural hydrophilic properties of cellulose maximize protein recovery by reducing protein adsorption. In addition, cellulose membrane can maintain high LRV value and filtration flow rate in long time filtration. Moreover, it has broad applicability and is suitable for virus removal process of multiple biological products.

Valpha® PES uses modified polyethersulfone material, and its unique asymmetric structure provides more possibilities for high process flux. At the same time, tofflon unique modification and membrane preparation process provides Valpha PES membrane very low protein adsorption and high flow rate. It really achieves the target of high flux, high load and high flow rate. It is truly characterized by high throughput, high loading and high retention.

Requirements for virus clearance

- ✓ Effective retention
 ✓ High yield
- ✓ Easy to use
- ✓ Shorter filtration time
- ✓ High capacity
- ✓ Can be validated
- ✓ Process robust







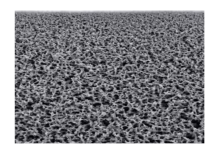
Meet your needs for virus removal

- Reliable virus removal ability
- Mice Minute Virus (MMV) reduction rate LRV≥4
- Antibody protein (150kD) recover rate ≥99%
- High efficient virus filtering unit
- Water based integrity testing, easy to test
- Single-use design, no cleaning validation required

- Sterility
- High pressure resistance
- High capability
- High flux, shorten the lot filtration time
- No additional filtration systems such as stainless steel clamps and accessories are required

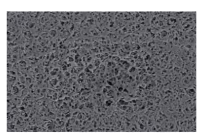
Unique virus retention membrane technology

Valpha® PES



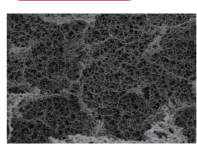
- High flux
- High retention
- Suitable for mab and clean drug luquid

Valpha® RC



- High retention
- High capacity
- Stable flux, suitable for polyclonal antibody, plasma products, biochemical extracts and other complex feed liquid

Valpha® RC HP



- High retention
- High capacity
- Designed for high concentration plasma products



Process development scale Syringe filter

- √Unique vent design
- √Effective membrane area of 2.8cm²
- √Suitable for process development, virus removal study and small scale process
- $\sqrt{100\%}$ in manufacture integrity test

Pilot scale 2.5"/5"/10" filter



- √ Suitable for pilot scale
- $\sqrt{\text{Pleated cartridge and capsule design}}$, easy to use.
- $\sqrt{}$ Gamma resistance (PES), Autoclavable and SIP tolerance (RC/RC HP)
- $\sqrt{100\%}$ in manufacture integrity test

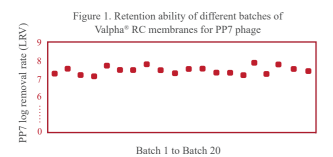
Process scale 20"/30" filter

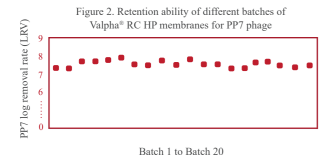


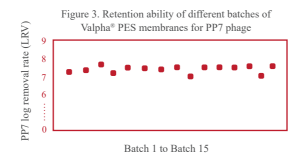
- √ Production scale
- $\sqrt{\text{Pleated cartridge and capsule design,}}$ easy to use.
- $\sqrt{}$ Gamma resistance (PES), Autoclavable and SIP tolerance (RC/RC HP)
- $\sqrt{100\%}$ in manufacture integrity test

Technical specification

PP7 phage retention capability







The photo shows the performance of Valpha® virus removal filter membrane on PP7 phage retention. It can be seen from the results that the three virus removal filter membranes, Valpha® PES, Valpha® RC and RC HP, have a strong virus removal performance as evidenced by the fact that for PP7 phage LRV≥7.

Valpha® Virus Removal Filter

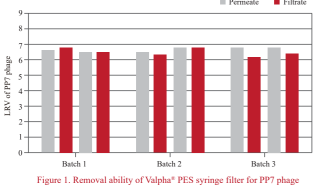


Product parameters

Membrane material			Pl	ES					RC					RC HP		
Specification(inch)	1	2.5	5	10	20	30	2.5	5	10	20	30	2.5	5	10	20	30
Effective filtration area(m²)	0.04	0.12	0.25	0.5	1	1.5	0.16	0.4	0.8	1.6	2.4	0.16	0.4	0.8	1.6	2.4
Component form			Caj	osule						C	Capsule,	Cartridg	e			
Connection		TC connector at both ends(Capsule); 226(Cartridge)														
Material		PP(Polypropylene)														
ISO*9001 quality standard		The R&D, production and delivery process of the products follows the requirements of quality management system of ISO9001:2015														
Membrane PP7 retention testing		Valpha® membrane - Phage PP7 retention test result indicates thatat the challenge level of 10 ⁷ pfu/cm ² , LRV> 4.0														
Device PP7 retention testing		Valpha® filter - Phage PP7 retention test result indicates that at the challenge level of 10 ⁷ pfu/cm ² , LRV> 4.0														
Fiber releasing				In c	omplia	nce with	the non	e fiber r	elease st	tandard	of 21CI	FR 21.3	(b)(6)			
USP <87> cytotoxicity test		The	compen	ent of Va	ılpha® f	lter hav	ve been to USP<		meet the		rds of c	lass VI p	olastics(accordin	g to	
USP <88> biological reactivity		The	compens	ent of Va	lpha® fi		re been to SP<88>					lass VI p	lastics(accordin	g to	
Sterilization method		Gan	nma resi	stance					Autocl	avable(0	Capsule); SIP(0	Cartridg	ge)		
Maximum operation pressure		Forward: 4.1bar(60psi)@4~30°C														
Bacterial endotoxin		Acco	rding to	Chinese			a 2020 e otoxin co						ed for e	ndotoxii	n and	

Technical index

PP7 phage retention performance data



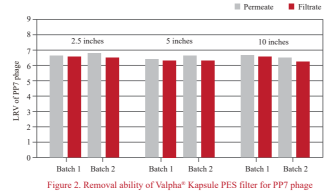
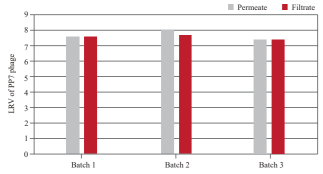


Figure 1. Removal ability of Valpha® PES syringe filter for PP7 phage



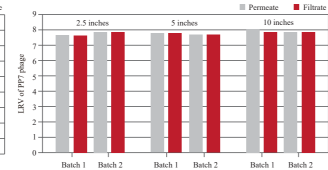
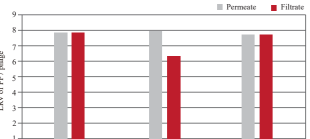


Figure 3. Removal ability of Valpha® RC syringe filter for PP7 phage



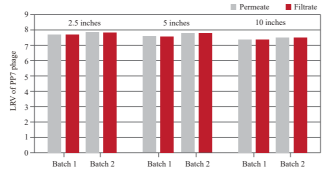


Figure 4. Removal ability of Valpha® Kapsule RC filter for PP7 phage

Figure 5. Removal ability of Valpha® RC HP syringe filter for PP7 phage

Figure 6. Removal ability of Valpha® Kapsule RC HP filter for PP7 phage

Water flux test data

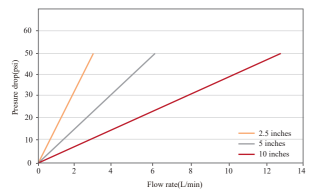


Figure 1. Valpha® Kapsule PES filter - Pressure drop and liquid flow rate

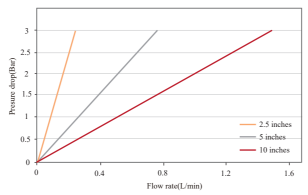


Figure 2. Valpha® Kapsule RC filter - Pressure drop and liquid flow rate

Water flux test data

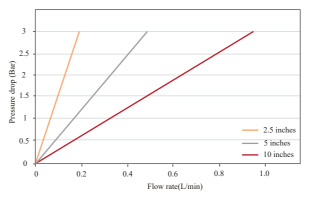


Figure 3. Valpha® Kapsule RC HP filter - Pressure drop and liquid

Order information

Valpha® Syringe filter





Product Type:

Product Scale:

D S 2 Membrane Area:

V1=Virus syringe

VR=Valpha RC RH=Valpha RC HP VP=Valpha PES

PD=Process development scale

S28=2.8cm²

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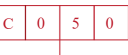
Valpha® Capsule filter







 \mathbf{C} 0 5





Product Code: V=Valpha

Component form: K=Kapsule

Membrane P=PES R=RC

H=RC HP

Specification: F005=1 inch

A012=2.5 inches B025=5 inches

C050=10 inches D100=20 inches E150=30 inches

Connector type:

SH=inlet TC25, outlet 1/2 to 1/4 HB(1 inch)

HH=1/2 to 1/4HB(1 inch) SS=TC25, In line(1/2.5/5 inches)

TT=TC50, In line(10/20/30 inches) LL=TC50, T line(10/20/30 inches)

Valpha® Cartridge filter













Product Code: V=Valpha

form: C=Cartridge

Component

R=RC H=RC HP

Membrane

Specification: A012=2.5 inches B025=5 inches C050=10 inches D100=20 inches

E150=30 inches

Connector type: 7=226

Vmax Equipment Set For Virus Filter

Filtration Experimental Equipment Set (Model:VHPDKIT)

Support for process development



Parts list

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Number	Product description	Quantity	Picture
1	Iron support	1	
	Relief valve (with pressure gauge)	1	~ ~
2	No.6 trachea	3	
	Gas valve switch	1	
	Diameter 64 fast-fit clamp (TC64)	2	
	Silicone gasket	2	b
3	Three-way valve	1	
3	NPT1/4 pagoda turn 64 quick connecter	1	
	Two-way luer connector	1	
	Translucent storage tank	1	

NOTE: Syringe virus removal filter, balance and collection bottles are not included in the set.

Vshell® Virus Pre-filter

Effectively improve the capacity of the virus pre-filter

Virus filteration step is often resulting in low filtration capacity, increased cost, increased process time and other problems due to various reasons. Without changing the state of the sample, adding a pre-filter is the most effective solution increase the capacity.

Vshell® Pre-filter is specially designed by Tofflon which can work together with Valpha Virus retention filter to provide a reliable virus safety garantee for the virus fitration process.

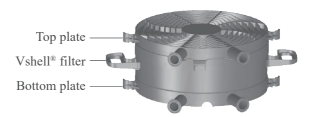
Selection of virus pre-filter

The purpose of virus pre-filter is to remove various impurities or contaminants such as protein aggregates, DNA, and other trace substances. The size of the pre-filter also has a significant impact on the viral filter capacity.

Tofflon provides a variety of specifications of pre-filter filters, according to different process requirements, to achieve robust virus removal purposes.

Product features

- Easy-to-use
- High flow rate, high capacity, high flow
- Improve the economic efficiency of virus removal filtration



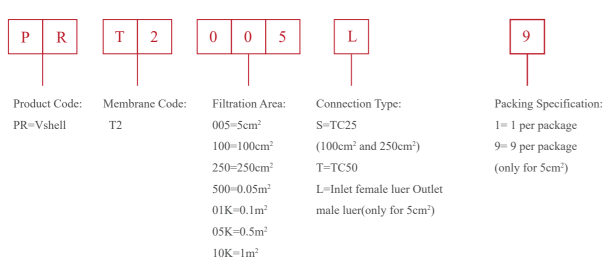


Product parameters

Spe	cification	5cm ²	100cm ²	250cm ²	500cm ²	1000cm ²	$0.5m^2$	$1m^2$			
Picture		÷			M D			D			
	Туре	Syringe filter	Capsule filter	Capsule filter	Capsule filter	Capsule filter	Plate filter	Plate filter			
Size	Diameter(mm)	36	94	94	94	94	442	442			
	Height(mm)	16.2	143	203	331	578	106	147			
Co	nnection	Luer	TC	25		TC	C50				
Ap	plication	R&D	R&D	Pilot scale	Pilot scale	Pilot scale	Process Scale	Process Scale			
C	apacity	0-1L	1-20L	1-20L	1-100L	1-100L	100-500L	100-2000L			
Material	Membrane	Diatomite, cellulose, perlite, positive charge resin									
Material	Housing			P	olypropylene						
Pre	ssure test	25°C water bath, maintain 50 psi for 24 hours, filter no gas leakage 40°C water bath, maintain 40 psi for 24 hours, filter no gas leakage									
Hydrau	lic stress test			40psi,	30min, 4cyc	cles					
	TOC		A	After flushing 1	00 L/m^2 , TOC	$was \leq 4mg/L$					
Con	ductivity	After flushing 100 L/m² , conductivity was 10 μ S/cm									
Bacteria	al endotoxins		After	flushing 100 L	m ² , endotoxin	was < 0.25 EU	J/ml				
Bios	safety test	The ma	aterial used me	eet the requiren	nents of USP <	87> and USP	<88> biosafety	y test			

Vshell® Holder and Accessories: See "Depth Filter Holder and Accessories".

Ordering information



Felix 6KP Integrity Tester

Product features

- Portable, easy to use, automated
- Complete integrity test of filter and process equipment accurately and reliably
- Simple and intuitive user experience
- Flexible custom optional Settings



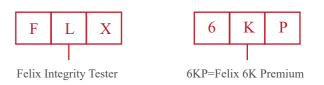
Product parameters

Tech	nnical index	6KP		
Inp	out voltage	(100-240)V AC		
Inpu	t frequency	50/60Hz		
In	put power	160W		
A	udit trail	Y		
Scre	en material	Resistive screen		
	mensions W × H /mm)	360×280×280		
	Diffusion flow	(0.1—600)ml/min		
Detection	Water intrusion	(0.01—100)ml/min		
range	Bubble point	(100—8000)mbar		
	Pressure maintenance	(100—8000)mbar		
	Diffusion flow	0.01ml/min		
Sensitivity	Water intrusion	0.01ml/min		
Schsitivity	Bubble point	50mbar		
	Pressure maintenance	1mbar		
	Diffusion flow	±4% or 0.1ml/min		
Precision	Water intrusion	±4% or 0.02ml/min		
Trecision	Bubble point	±50mbar		
	Pressure maintenance	1mbar		

Product compliance

- Support multi-level user access Settings
- Ensure data integrity
- Audit trail function

Ordering information





TanFlux® Series TFF Membrane

In order to meet the needs of different applications, we offer two defferent kinds of membrane:

Modified PES

Polyethersulfone



PES membrane can tolerate the irritant chemical substances used in cleaning, decontamination and disinfection. The modified PES membrane improves on the basis of the traditional PES membrane to reduce nonspecific protein binding. The technique also makes the modified membrane tougher for withstanding process disturbances and extreme operating conditions. High flux and retention performance make processing faster and ensure higher yield, thus reducing the processing time.

Composite Flawless RC Regenerated cellulose



The high anti-fouling and anti-blocking performance and low protein adsorption of compositeflawless RC membrane ensure excellent product retention, recovery and higher yield. Flawless RC membrane is a flawless membrane made with RC membrane on microporous substrate. Compared with traditional membrane, it has excellent toughness. The mechanically robust design provided by composite material technology enables it to withstand process disturbances and extreme operating conditions.

Product parameters

Membrane Material	Modified PES(polyethersulf	fone)	CompositeFlawless RC (regenerated cellulose)
Filtration Type	Ultrafiltration	Microfiltration	Ultrafiltration
Molecular weight cutoff	5/8/10/20/30/50/70/100/300/500KD	0.1/0.22/0.45µm	1/2/3/5/10/30/100/300KD
Relative flow rate	Fast	Medium	
Protein adsorption	Low		Very low
pH range	1-14	2-13	
Characteristics	High flux, good chemical comp	Very low protein adsorption, hard to block, resistant to organic solvents	

Quality assurance

The same equipment, process and quality standards are adopted for all TanFlux® TFF membrane cassettes in production. 100% integrity test and visual inspection are conducted for each manufacturing batch of membrane cassettes in production to ensure that each membrane cassette is integral, intact in appearance and is up to the delivery quality standards. Each membrane cassette has a unique serial number in the batch number and a separate quality certificate.

The consistent performance of membrane's material guaranteed in the automatic manufacturing process ensures the consistency and high repeatability from scale expansion to scale reduction and between batches. All TanFlux® membrane cassettes are produced under the quality system that that refer to GMP requirements.

TanFlux® II TFF Cassette



TanFlux® II TFF Cassette are produced automatically to ensure the consistency of performance and linear scalability among the cassettes of different lots. The simplified design enables users to install and operate quickly and easily.

With excellent chemical compatibility, TanFlux® II TFF Cassette can be compatible with various chemical cleaning agents to ensure safe and effective cleaning and disinfection of the TFF system.

Product features

- From membrane to finished cassettes fully indepentent and localized.
- Robust and reliable design of the housing which can withstand high pressures and temperature.
- Easy to install and clean to ensure the reliability of the yield
- Multiple flow channel options
- Fast and reliable linear scale-up from lab to production scale
- Exllent chemical compatibility
- Extrem Low E&L level
- High separation efficiency

Process upgrading

1 Thermal welding sealing process

The unique thermal welding sealing process replaces the traditional glue injection process. Significantly improve the alkali resistance and chemical compatibility.

2 New resin formulation

The molding process adopts a new resin formula, while retaining the traditional silicone packaging process, which has higher hardness and better process stability.

3 More channel options

On the basis of the existing D channel, the new V (open) channel is compatible with different application scenarios.

4 More uniform pore size distribution

The optimized membrane process makes the membrane pore size distribution more uniform, the sieving effect is strong, and the protein separation efficiency is higher.

Reliable scale up from R&D to production

TanFlux® II has four sizes: 80 cm², 0.1 m², 0.5 m² and 2.5 m². All TanFlux® II TFF Cassette membrane are made from the same materials and have the same channel structure, length and height, which enables TanFlux® II TFF Cassette membrane of various specifications to maintain the same performance, and truly achieve scale up from 100ml to thousands of liters. TanFlux® II TFF Cassette membrane is designed with a strong resin and silicone housing to protect the membrane's surface against the impact and potential damage.

Product specification

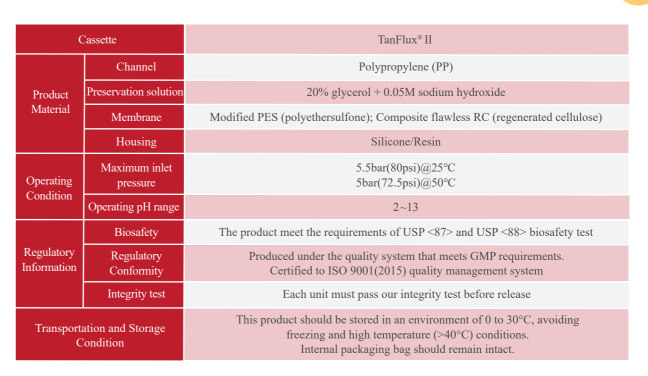
Membrane area	80cm ²	$0.1 m^2$	$0.5 {\rm m}^2$	2.5m ²
Resin housing		Star Start Francisco	Office and a second	Space Terfor 1 Course Ort tomoget President Ples to Act Ser Contractor do
Silicone housing		See North State States	Barrer to be a before the second	SAP Terful® Cessels Of PROCES Filtration reside Code Ufficientes 9
Procesing volume	100mL-1L	200mL-10L	2L or large-scale production	10L or large-scale production
Cassette length (mm)	207	207	205	205
Cassette width (mm)	56	56	178	178
Cassette thickness (mm)	4	15-16	15-16	70-75

Silicone housing

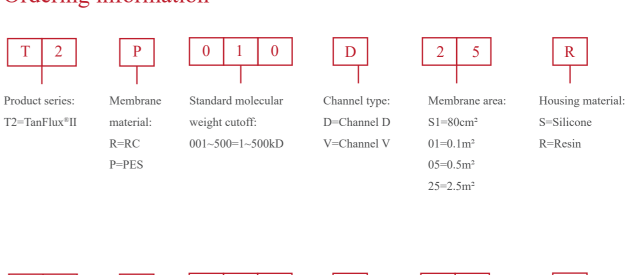
Resin housing



Product parameters



Ordering information



				20 2.0.11	
T 2	P	M 2 2	D	2 5	R
Product series:	Membrane	Standard molecular	Channel type:	Membrane area:	Housing material:
T2=TanFlux®II	material:	weight cutoff:	D=Channel D	S1=80cm ²	S=Silicone
	P=PES	M22=0.22um		$01=0.1 \text{m}^2$	R=Resin
		M45=0.45um		$05=0.5m^2$	
		M10=0.10um		25=2.5m ²	

TFF Holder & Accessory

Suitable for different specifications of membrane cassette, to meet different process requirements

Product features

- The holder body is made of stainless steel 316L
- The structural design complies with the cGMP medical grade standard. The diaphragm valve and pressure gauge with sanitary dead corners
- Stable filtration effect

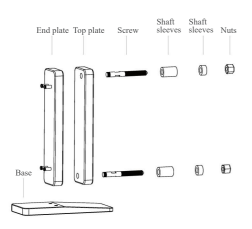
Mini TFF Holder

TFH100 (Including accessories)

Max. Holding Capacity: 80cm² cassttes*5

The minimum working volume of the mini TFF holder can reach 20mL

Holder assembly diagram





Accessories information

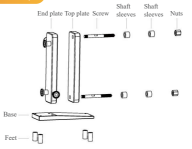
Product picture	0					net miles to
Product series	Pressure tap	Sanitary grade 0-0.6 MPa pressure gauge	10-32UNF threaded to Luer connection	1/8 "Luer male connection	Luer Plug	Two-way value or Luer connection
Quantity	2	2	4	3	1	2

0.1m² Stainless Steel Holder

TFH01 (Including accessories)

Max. Holding Capacity: 0.1m² cassttes *3

Holder assembly diagram





Accessories information

Product picture			00	00	9	0				
Product series	φ9.8mm three-way pipe	Sanitary grade 0-0.6 MPa pressure gauge	TC25 clamp	TC50 clamp	φ8-12mm hose clamp	0.5 " clamp gasket		12.7*1.5mm manual diaphragm valve	φ25.4mm nausea board	φ6-25mm hose connection
Quantity	2	2	11	2	5	11	2	2	1	3

0.5m² Stainless Steel Holder

TFH05 (Including accessories)

Max. Holding Capacity:

0.5m² cassttes *5,

2.5m² cassttes *1
(Optional extension screw)

Holder assembly diagram

Shaft sleeves

Shaft sleeves

Shaft sleeves

Handle

Top plate

Bottom plate



Accessories information

Product picture			00	00	9	0			3	
Product series	φ9.8mm three-way pipe	Sanitary grade 0-0.6 MPa pressure gauge	TC25 clamp	TC50 clamp	φ8-12mm hose clamp	0.5 " clamp gasket		12.7*1.5mm manual diaphragm valve	φ25.4mm nausea board	φ6-25mm hose connection
Quantity	2	2	11	2	5	11	2	2	1	3

SaiClear® Depth Filter

Multiple grades of depth filters provide suitable solutions for different application scenarios.

Used in clarification and downstream filtration processes

Depth Filter is commonly used for pre-filtration, such as fermentation fluid clarification, removal of cell debris and DNA, preconditioning in plasma products and other biological products.

SaiClear® Depth Filter, developed for demanding clarification applications in the biotechnology and pharmaceutical industries, significantly reduce the difficulty of downstream purification.

Product features

- With diatomaceous earth and cellulose as membrane materials, it has a high contaminant capacity
- Higher impurity removal capacity through mechanical interception and electrostatic adsorption
- Combine multiple clarification steps into one device to optimize the clarification process
- A full line of products is available for process reduction and scale-up

Filter application

- Clarification filtration of fermentation and cell culture fluids
- Filtration of plasma products
- Chemical medicine impurity removal, decarbonization filtration
- Remove particles and glue from process intermediates

Clarification filtration

The purpose of clarification is to effectively separate the cells, cell fragments and other colloidal substances in the harvest solution to provide an impurity-free feed solution for downstream purification. Clarification of cell harvest solution is the first step in middle and downstream purification of biopharmaceutical processes, usually using a combination of centrifugation, tangential flow filtration, flocculation, and depth filtration. Depth filters technology, because it can improve product quality, simplify operation, and more and more respected by the industry. The harvested clarification filtration usually uses two grades of depth filters: the first grade uses a filter medium with a large aperture to remove cells or cell debris; The second grade uses a filter medium with a tight aperture to remove colloidal impurities.

Depth Filter

18

SaiClear® P Series and S Series depth filters have been developed to meet different process requirements. The filter medium inside the filter is the same as the filter medium in the traditional depth filter system, and there are many choices of accuracy and models. In addition, SaiClear® has a small footprint, saving clean room space. Compared to traditional depth filtration systems, the SaiClear® installation position is flexible and easy to operate for process development.

Product picture

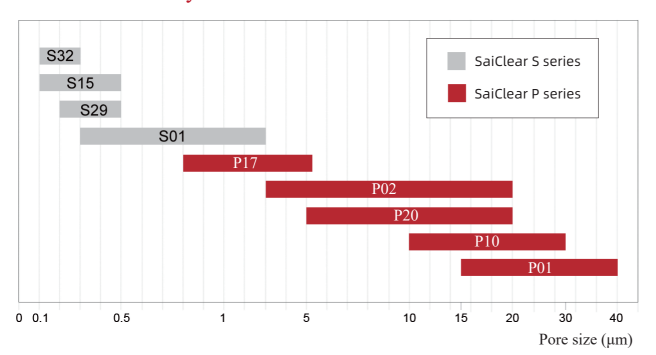
23cm ²	100cm ²	250cm ²	480cm ²	1000cm ²	$0.45m^{2}$	0.9 m 2
		Manus Sylven	H		Service Servic	

Product parameters

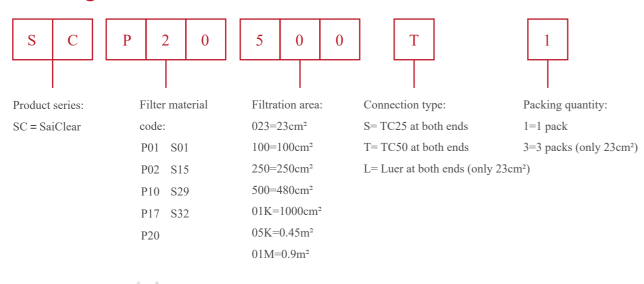
From process development to mass production, we provide you with the right solution

Specif	ication	23cm ²	100cm ²	250cm ²	480cm ²	1000cm ²	$0.45m^{2}$	0.9m ²			
Filter	Filter type		Capsule	Capsule	Capsule	Capsule	Plate	Plate			
Equipment	Diameter mm	63.8	94	94	94	94	442	442			
size	Height mm	51	143	203	331	578	106	147			
Conn	ection	Luer	TC	225		TC	250				
Appli	cation	Ra	&D	Pilot	scale		Production				
Processin	g volume	0-	5L	1-2	20L		20-2000L				
Material	Membrane material			Diatoma	aceous earth,	cellulose					
iviateriar	Housing	Polypropylene									
Integr	ity test	No gas leakage, aerosol permeability meets the requirements									
Pressi	ıre test	25°C water bath, maintain 50psi pressure for 24 hours, filter no gas leakage 40°C water bath, maintain 40psi pressure for 24 hours, filter no gas leakage									
Hydraulic	stress test	Rinse the filter at 40psi for 30 minutes and repeat this cycle four times before the filter passes the integrity test									
TO	OC .	After rinsing	g with 100L/n	n² volume of p	urified water,	sample the fi	ltrate. TOC va	lue ≤6mg/L			
Condu	ıctivity	After rinsing with 100L/m² volume of purified water, sample the filtrate. Conductivity value ≤15µS/cm									
Bacterial	endotoxin	The content of endotoxin in the filtrate was less than 0.25EU/mL									
Biosafet	ty testing	Tl	ne materials us	sed meet USP	<87> and US	P <88> biosaf	ety requireme	nts			

Filtration accuracy



Ordering information



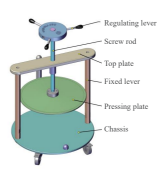


Holder and Accessories

Various specifications of depth holder, suitable for different specifications of filter packaging, to meet different process requirements

Product features

Compared to similar products on the market, the provided holder can precisely adjust the pressure level.





Product parameters

Specification	Pilot scale holder	Production grade holder
Product Picture		
Length*Width*Height(mm)	580*530*1000mm	775*575*1740mm
Weight	733kg	178kg
Max. Holding Capacity	0.9m ² *2	0.9m ² *9
Surface finishment	ra3.2	ra3.2
Adjustable torque	5-35N·m	5-35N·m

Ordering information

SCFH05M	Pilot scale holder
SCFH10M	Production grade holder

Item	Product name	Specification	Unit
SCCP015TT	Saiclear upper cover	0.45m ² , 0.9m ² general	piece
SCCP015BT	Saiclear lower cover	0.45m ² , 0.9m ² general	piece
SCCP015MT	Saiclear Partition	0.45m ² , 0.9m ² general	piece

SaiPress® Sterilizing-grade Hydrophilic Filter

High flow rate sterilizing-grade polyether sulfone (PES) filter

Product features

- Sterilizing-grade polyether sulfone (PES) material, can withstand multiple autoclaves
- Higher flow rates at the same level, thus reducing the process footprint and maximizing benefits
- Ready-to-use models for gamma irradiation
- Good chemical compatibility for pH1-14
- A full line of products is available for process reduction and scale-up

Filter application

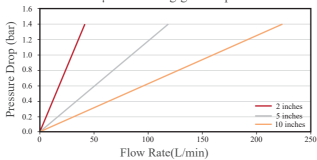
- Buffer, pH regulator filtration
- Process additives and intermediates filtration
- Chemical synthetic medium filtration
- Water-based feed filtration

Filter type

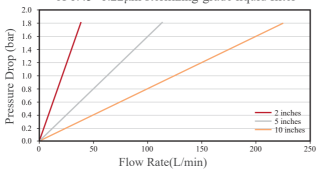
- Ready-to-use capsule filters
- Cartridge filter
- Capsule filters sterilized by gamma irradiation
- Disk filters and customized filters

Water flux test data

Relationship between pressure drop and flow rate of 0.22µm sterilizing-grade liquid filter

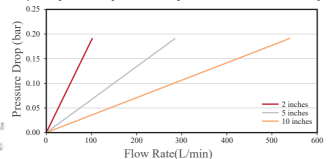


Relationship between pressure drop and flow rate of $0.45 + 0.22 \mu m$ sterilizing-grade liquid filter





0.45µm liquid filter pressure drop and flow rate relationship



Product parameters

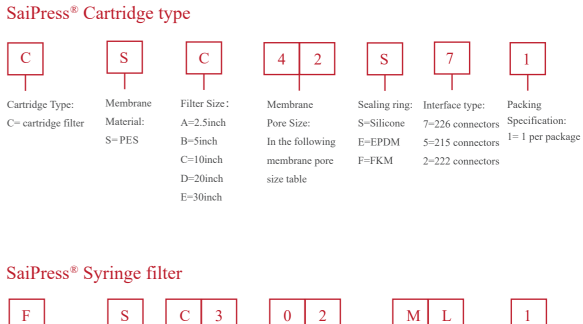
			Disk filter Capsule filter Cartridge filter											
	Filter type		Disk filter			Capsu	le filter				C	artridge fil	ter	
No	minal size (inch	•	Ф50mm	1	2.5	5	10	20	30	2.5	5	10	20	30
	Effective filtration Single layer			380cm ²	0.16	0.33	0.66	1.32	1.98	0.16	0.33	0.66	1.32	1.98
are	ea (m²)	Double layer	13.8cm ²	350cm ²	0.14	0.28	0.57	1.14	1.71	0.14	0.28	0.57	1.14	1.71
Filter	membrane pore	size		$0.1\mu\text{m}/0.22\mu\text{m}/0.45\mu\text{m}/0.65\mu\text{m}/0.8\mu\text{m}/1.2\mu\text{m}/0.22+0.1\mu\text{m}/0.45+0.1\mu\text{m}/0.22+0.22\mu\text{m}$										
Titter	memorane pore	SIZC	$0.45 + 0.22 \mu m/0.65 + 0.22 \mu m/0.8 + 0.22 \mu m/1.2 + 0.22 \mu m/0.8 + 0.45 \mu m/1.2 + 0.45 \mu m/0.8 + 0.45 \mu m/0.$											
	Filter men	ıbrane		Hydrophilic polyether sulfone (PES)										
	Supporting layer structural cor	er/Housing	Polypropylene (PP)											
Material	Vent O-r	ing				S	ilicone							
	O-rin	g									Silicor	ne/EPDM/	FKM	
					Г	1. 5.51	(00 ')	@350G+		т	Zomrrond .	5 5hau(90u	:\@25°C	
Ma	ximum operatin	g			Forw	ard: 5.5b	ar(80psi) ar(25psi)	_		Ι	orward.		osi)@25°C (psi)@80°C	
pre	essure difference					1.70	ur(25psi)	w,00 C				1./bai(23	ps1)@60 C	_
					Reve	rse: 2.1b	ar(30psi)(@25°C			Reverse:	2.1bar(30	psi)@25°C	
Bubble po	int (100% IPA (@ 25°C)						0.1μm:≥25	600mbar					
Bubble po	int (pure water (@ 25°C)			0.2	2μm:≥340	0mbar	0.45μm:≥	2000mbar	0.65μπ	n:≥1500ml	bar		
•	r diffusion flow er @ 25°C@280			≤2.5	≤7.5	≤15	≤30	≤60	≤90	≤7.5	≤15	≤30	≤60	≤90
•	er diffusion flow er @ 25°C@280			≤2.5	≤7.5	≤15	≤30	≤60	≤90	≤7.5	≤15	≤30	≤60	≤90
			Gamma not applicable	SIP not applicable							Gam	ıma not ap	plicable	
Ste	rilization metho	d	Offline: 126°C / 60 minutes, 5 times	Code R: Tolerance 40KGy gamma irradiation, autoclave not allowed Code A: Offline: 126°C/60min, 3 times, Gamma irradiation not allowed						(In t	SIP :126 he steriliza downstrear	°C / 60mir ation proce	ss, the ups difference	tream
	Pressure test			No lea	akage dete	cted under	0.65MPa	pressure						
TOC	and conductivi	ty	Accordin China Phar according t	macopoeia	a 2020 Edi	tion (0682) and the		on method	of electric	cal conduc	tivity of ph	armaceutio	eal water
	Cleanliness			Mee	ts the "no	fiber relea	se" filter	standard as	s defined in	n 21 FR21	0.3 (b) (6))		
Ba	cterial endotoxi	n	Meets the "no fiber release" filter standard as defined in 21 FR210.3 (b) (6) According to the bacterial endotoxin test method stipulated in the General Rule of China Pharmacopoeia 2020 Edition (1143), the endotoxin content of the filter solution was determined, and the content of endotoxin in the filter solution was lower than 0.25EU/ml											
В	iosafety testing	All mater	rials of the	filter cart	_		SP < 88 >, i		-	lluation of	ClassVI p	lastic prod	ucts	
Ва	cterial retention	The test					eudomonas hat the min					etention of	this	
Regu	ılatory compliar	ıce		Pı	roduct pro	duction an	d quality	manageme	ent in acco	rdance wit	th ISO: 90	01 (2015)		
	Shelf life							3 yea	rs					
Storag	e and transporta						ent of 0 to 3							

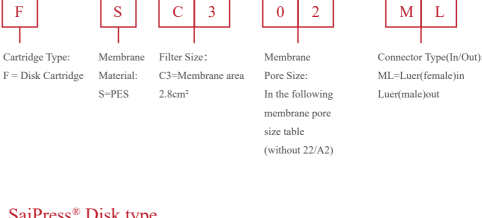
Ordering information

SaiPress® Capsule

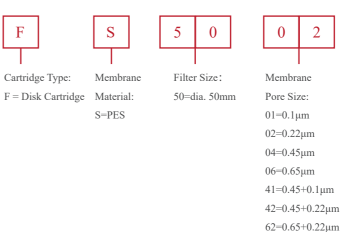


Ordering information





SaiPress® Disk type





HH= 1/2 to 1/4 hose





Packing

Specification:

1= 1 per package

9= 9 per package

Packing Specification: 1= 1 per package

Membrane	Single layer	01=0.1μm/02=0.22μm/04=0.45μm/06=0.65μm/08=0.8μm/12=1.2μm
pore size	Double layer	$21 = 0.22 + 0.1 \mu m/41 = 0.45 + 0.1 \mu m/22 = 0.22 + 0.22 \mu m/42 = 0.45 + 0.22 \mu m/62 = 0.65 + 0.22 \mu m/82 = 0.8 + 0.22 \mu m/A2 = 1.2 + 0.22 \mu m/84 = 0.8 + 0.45 \mu m/A4 = 1.2 + 0.45 \mu m/A$

SaiPress® HF Sterilizing-grade Hydrophilic Filter

High flux sterilizing-grade hydrophilic polyether sulfone (PES) filter membrane

Product features



The flow rate is more than 2.5 times that of the same type of sterilizing-grade filter membrane



High capacity, the same filtration area, larger processing capacity



Good chemical compatibility, suitable for various solutions of pH 1-14, including NaOH solutions



Process improvements increase efficiency and reduce costs

Filter application

• Buffer, pH regulator filtration



Product parameters

	1														
	Filter type		Syringe filter			Capsul	e filter				С	artridge fil	lter		
No	ominal size (i	nch)		1	2.5	5	10	20	30	2.5	5	10	20	30	
Effective	filtration	Single layer	2.8cm ²	380cm ²	0.16	0.33	0.65	1.3	1.95	0.16	0.33	0.65	1.3	1.95	
area	(m ²)	Double layer	2.8cm ²	350cm ²	0.14	0.28	0.55	1.1	1.65	0.14	0.28	0.55	1.1	1.65	
Filter	membrane p	ore size		0.22μm/0.45μm/0.22+0.22μm/0.45+0.22μm/0.8+0.45μm											
	Filter m	embrane	Hydrophilic polyether sulfone (PES)												
		ing layer					I	olypropyle	ene (PP)						
Material	Housing compon	/structural ents		Polypropylene (PP)											
	Vent (O-ring				Si	licone								
	О-1	ing									Silic	one/EPDN	1/FKM		
	aximum oper					1.7ba	ar(25psi)@	80°C			orward: 5 1 everse: 2.	.7bar(25ps	si)@80°C		
Bubble po	oint (100%IF	PA@25°C)	0.22μm: ≥3400mbar												
	er diffusion f ter @ 25°C@	low (ml/min) 2800mbar)		≤2.5	≤7.5	≤15	≤30	≤60	≤90	≤7.5	≤15	≤30	≤60	≤90	
						SIP not	applicable				Gamm	a not appl	icable		
Ste	erilization me	ethod		Code R: Tolerance 40KGy gamma irradiation, autoclave not allowed Code A: Offline: 126°C/60min, 3 times, Gamma irradiation not allowed							Offline:126°C / 60min, 25 times SIP:126°C / 60min, 25 times (In the sterilization process, the upstream and downstream pressure difference of the cartridge is ≤300mbar)				
	Pressure tes	t		No lea	kage detec	cted under	0.65MPa w	ithin 24h							
ТОС	C and conduc	ctivity	China Ph	ing to the dearmacopoeing to the gene	a 2020 Ed	ition (0682) and the d	eterminatio	n method o	f electrical	conductiv	ity of phar	maceutica	l water	
	Cleanliness				Meets	the "no fib	er release"	filter stand	ard as defin	ed in 21 F	R210.3 (b)	(6)			
Ba	ecterial endot	oxin		ccording to on (1143), t			of the filte	r solution v		ined, and th					
E	Biosafety test	ing	All ma	terials of th	e filter car	tridge mee	t the biosaf	ety assessn	nent of USI	o < 88 >, C	lassVI-70°	C grade pl	lastic mater	rials	
Ва	acterial reten	tion	The	test method					s (ATCC19				ention of th	nis	
Reg	ulatory comp	liance			Product p	roduction a	and quality	manageme	ent inaccord	ance with	ISO:9001	(2015)			
	Shelf life							3 yea	rs						
Storag	ge and transp	ortation	Th	is product s					30°C, avoid)	
In	soluble parti	cles	Unless otherwise specified, the number of particles containing 10µm and more than 10µm in each 1ml filter rinse solution shall not exceed 25 particles, and the number of particles containing 25µm and more than 25µm shall not exceed 3 particles												

Ordering information





Filter type:

C= cartridge filter



H=HF



Filter Size:

A=2.5inch

B=5inch

C=10inch

D=20inch

E=30inch



Membrane

02=0.22μm

 $04=0.45 \mu m$

22=0.22+0.22µm

42=0.45+0.22μm

84=0.8+0.45µm



S=Silicone

E=EPDM

F=FKM





7=226 connector

Specification: 1= 1 per package



Membrane

Material:

H=HF



Filter Size:

 2.8cm^2

C3=Membrane area

Membrane

Pore Size: 02=0.22μm 04=0.45μm

22=0.22+0.22μm

 $42=0.45+0.22 \mu m$

Luer(male)out

Connector Type(In/Out):

ML=Luer(female)in

Packing Specification:

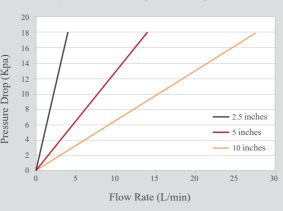
> 1= 1 per package 9= 9 per package

Test data

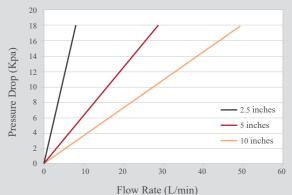
Filter type:

F=Disk filter

0.45+0.22μm SaiPress® HF pressure drop and flow rate diagram



0.22μm SaiPress® HF pressure drop and flow rate diagram





SaiPress® HF Pro Sterilizing-grade Hydrophilic Filter

Ultra-high Flux Sterilizing-grade Polyether sulfone (PES) Filter

Product features



The flow rate is more than 2.5 times that of the same type of sterilizing-grade filter membrane



High capatity, the same filtration area, larger processing capacity



Higher retention rates



Low dissolution and precipitation to ensure filtrate purity



Low adsorption, reduce filtrate loss

Filter application

- Chemical synthetic media filtration
- Final sterilization filtration

Filter type

- Sterilized packaging(Gamma sterilization)
- Syringe filter



Product parameters

	Filter type		Syringe filter			Capsu	ıle filter				Cartrio	lge filter		
No	minal size (inc	h)		1	2.5	5	10	20	30	2.5	5	10	20	
	e filtration a (m²)	Double layer	2.8cm ²	350cm ²	0.14	0.28	0.57	1.14	1.71	0.14	0.28	0.57	1.14	
Filter	membrane por	e size					0.22+0.1µ	ım/0.45+0.	1μm/0.45+().22µm				
	Filter mer	nbrane					Hydropl	nilic polystl	nersulfone ((PES)				
	Supportin	ng layer	Polypropylene (PP)											
Material	Housing/st componen		Polypropylene (PP)											
	Vent O-	ring				Sili	cone							
	O-rin	g									Silicone / E	PDM / FKM		
	ximum operati					1.7ba	r(80psi)@2 r(25psi)@8 r(30psi)@2	80°C			ward: 5.5ba 1.7ba verse: 2.1ba	r(25psi)@80	°C	
					Rever	. 2.10di	.(50psi)@2	3 C		RC	verse. 2.10d	1(30ps1)@23		
Bubble poi	int (100%IPA	@25°C)		≥2500mbar (100%IPA@25°C)										
	r diffusion flow er @ 25°C@28	` ′		≤2.5	≤7.5	≤15	≤30	≤60	≤90	≤7.5	≤15	≤30	≤60	
				SIP not applicable Gamma not applicable										
Ste	rilization meth	od		C		autoclave i Off-line: 12	Gy gamma not allowed 26°C/60min ot allowed	Offline:135°C* 30min, 25 times SIP:135°C* 30min, 25 times (In the sterilization process, the upstream and downstream pressure difference of cartridge is ≤300mbar)						
	Pressure test			N	o leakage	detected ur	nder 0.65M	Pa within 2	24h		-	-		
TOC	and conductiv	vity	Pharmacop	oeia 2020 E	Edition (068	82) and the	determinat	ion method	of electrica	l conductivity	cording to the y of pharmace nductivity val	utical water a	according to	
	Cleanliness			Meets the "no fiber release" filter standard as defined in 21 FR210.3 (b) (6)										
Ba	cterial endotox	in		_			lter solution	n was deter		the content o	ina Pharmaco f endotoxin ir	^		
В	Biosafety testing	or S		All mater	rials of the	filter cartri		*	oiosafety as astic materia		USP < 87 >, U	JSP < 88 >,		
Ва	acterial retentio	n	The	test method			efective Ps	eudomonas	s (ATCC19		ed for bacteria s 10 ⁷ cfu/cm ²	al retention o	f this	
Regu	latory compliance Product production and quality management in accordance with ISO:9001 (2015)													
	Shelf life		3 years											
Storag	e and transport	ation	This product should be stored in an environment of 0 to 30°C, avoiding freezing and high temperature (>40°C) conditions. The internal bag should be kept intact to prevent external particles or moisture from entering											
In	soluble particle	es			^		^		articles con		n 10µm in ead			

Ordering information



(Sterilized packaging)

R=Gamma sterilizable

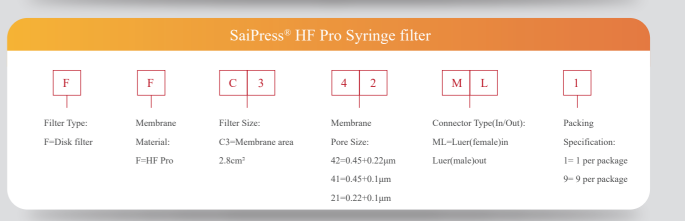
A=Autoclavable

down-out

right-out

T=Left-in and





Test data

1= 1 per package

0.45+0.22µm SaiPress® HF Pro pressure drop and flow rate diagram



F=HF Pro

E 30inch

42=0.45+0.22μm

41=0.45+0.1μm

21=0.22+0.1µm

SaiPress® PVDF Hydrophilic Sterile Filter

SaiPress® PVDF hydrophilic sterile filter is suitable for sterile filtration of various biopharmaceutical products. It is made of PVDF membrane with symmetric pore structure, uniform hydrophilicity, high porosity, and extremely low protein adsorption.

Product Features

- High porosity, extremely low protein adsorption
- Low extractables and good chemical compatibility for pH1-14
- Sterilizing-grade (0.22µm) PVDF material, can withstand multiple autoclaves
- Higher flow rates at the same level, thus reducing the process footprint and maximizing benefits
- 100% integrity testing ensures sterilization effectiveness

Product applications

- Filtration of colloidal
- Filtration of cell culture medium
- Filtration of High-viscosity liquid filtration and high-concentration liquid
- Filtration of sterile APIs and ophthalmic preparations
- Filtration of high-purity chemicals, intermediates, strong acid and oxidizing liquids
- Filtration of vaccines, biologics, blood products

Filter type

- Ready-to-use capsule filters
- Cartridge filter
- Capsule filters sterilized by gamma irradiation
- Customized filters

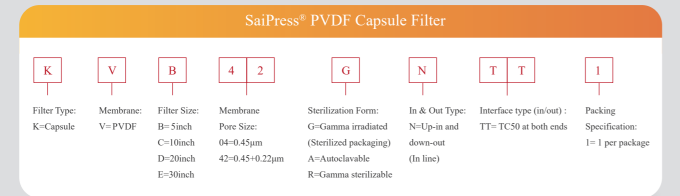


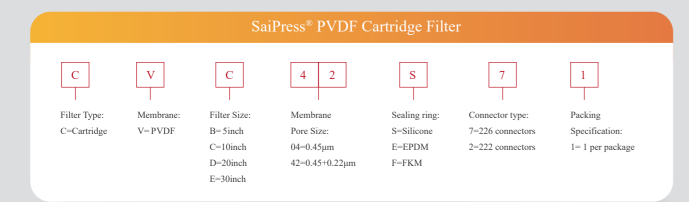


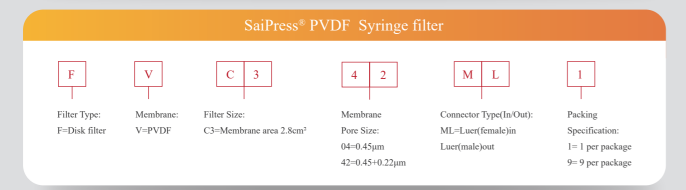
Product parameters

Filter type			Syringe Filter		Capsul	e Filter		Cartridge Filter					
		e (inch)		5	10	20	30	5	10	20	30		
	Effective filtration area (m²) Single layer Double layer		2.8cm ² 2.8cm ²	0.33	0.65	1.3	1.95 1.65	0.33	0.65	1.3	1.95		
	Filter membrane pore size		2.60111										
rinei ine		r membrane	0.45μm、0.45+0.22μm										
		orting layer	PVDF Polypropylene(PP)										
Material	Hous	ing/structural	Polypropylene(PP)										
		mponents ent O-ring			Silio	cone			-	-			
		O-ring			-	-			Silicone/El	PDM/FKM			
				Forw	ard: 5.5ba	ır(80psi)@2	25°C;	Forv	ward: 5.5ba	ar(80psi)@2	25°C		
		perating ference				r(25psi)@8				ar(25psi)@8			
press				Rev		ar(30psi)@2		Rev	erse: 2.1ba				
						(-1)				/ .			
	bble p	oint (a) 25°C)			0.45+0.22	μm:≥3400r	nbar 0.4	45μm:≥200	0mbar				
(pure	water (.u, 23 C)											
	5+0.22				40.6	25.0	27.0		40.6	25.0			
		ow (ml/min) C@2700mbar)		≤6.3	≤12.6	≤25.2	≤37.8	≤6.3	≤12.6	≤25.2	≤37.8		
).45µm sion flo	n ow (ml/min)		≤10	≤20	≤40	≤60	≤10	≤20	≤40	≤60		
		@1600mbar)											
			Offline:124°C/30min, 5times; STP.1249C/20i. 20vi										
Sterili	zation	method		134°C / 20min, 5times; SIP:124°C / 30min, 30tim 134°C / 20min, 5times; 134°C / 20min, 30tim									
			126°C / 60min, 5times;							ines,			
			According to the determination method of total organic carbon in pharmaceutical water according								-		
TOC ar	nd cond	ductivity	to the General Rule of China Pharmacopoeia 2020 Edition (0682) and the determination method of electrical conductivity of pharmaceutical water according to the general rule (0681), the detected										
									value is ≤ 1 .		actected		
C	leanlin	ess	N	leets the "	no fiber rel	ease" filter	standard as	defined in	21 FR210.3	3 (b) (6)			
									the General				
Bacterial endotoxin			Pharmacopo						ilter solutio wer than 0.2		mined,		
Disco-Catalana di an								ssment of U					
Biosafety testing					ClassVI gra								
Bacterial retention			0.45+0.22μm: bacterial reten										
Regulat	ory co	mpliance	Proc	duct produ	ction and q	uality mana	gement ina	ccordance	with ISO:90	001 (2015)			
S	Shelf li	fe					3 years						
Insol	uble pa	articles	Unless other 1ml filter rin		shall not e		articles, and	the number	er of particle				

Ordering information

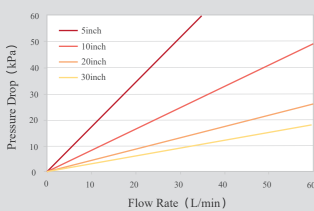


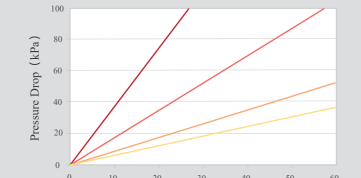




Test data

0.45µm SaiPress® PVDF pressure drop and flow rate diagram 0.45+0.22µm SaiPress® PVDF pressure drop and flow rate diagram





Flow Rate (L/min)

Filgiant® Sterilizing-grade Hydrophobic Filter

Ideal for gas and non-aqueous liquid filtration

As a sterilizing-grade gas filter composed of single-layer/double-layer polytetrafluoroethylene (PTFE) membrane and polypropylene components, the Filgiant® sterilizing-grade hydrophobic filter has good chemical compatibility and boasts its characteristics of high flow rate, high flux and no shedding of fibers, providing all kinds of filter sizes and connectors to meet your wide application needs.

Product features

- Made of the sterilizing-grade polytetrafluoroethylene (PTFE) materials, it can withstand multiple sterilization cycles;
- $\bullet~$ By physical retentionand charge adsorption, it can retent particles of size down to 0.01 $\mu m;$
- High-temperature and high-pressure tolerance;
- With good chemical compatibility, pH1-14 tolerance;
- Provide all scales products to meet your needs from R&D to manufacture.

Filter application

- Vent filter for storage tanks, fermentation tanks/WFI tanks;
- Gas sterile filtration of compressed air/clean pipelines;
- Input/output gases for bioreactors;
- Sterile filtration of non-aqueous liquids

Filter types

- Ready-to-use capsule filter
- Cartridge filter
- Disk filter and customized filters

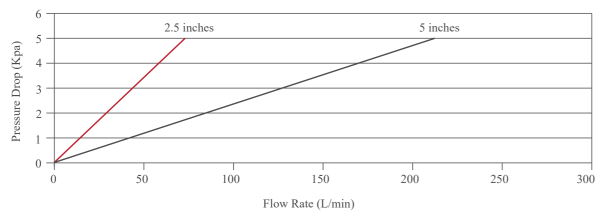


Connector types

TC25	TC50	1/4-1/2HB	215 Connector	internal insert Connector	222 Connector	226 Connector
	3			=		

Pressure drop and gas flowrate



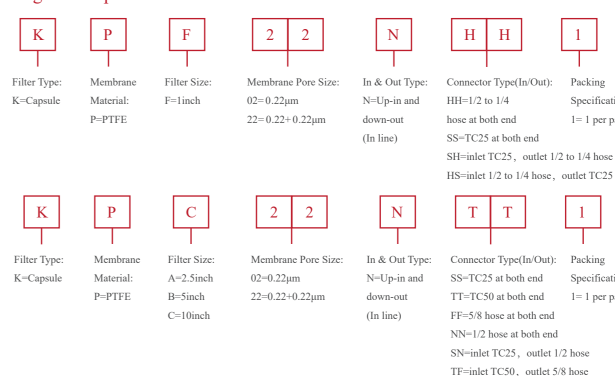


Product parameters

	Filter type	Disk filter	Capsule filter	Cartridge filter						
	Ther type		Single layer/	Double layer						
1	Nominal size	Ф50mm	1/2.5/5/10/20/30 inches	2.5/5/10/20/30 inches						
Men	nbrane pore size		0.22μm/0.22+0.22μm							
Effect	tive filtration area	13.8cm ²	0.04/0.16/0.33/0.65/1.3/1.95m ²	0.16/0.33/0.65/1.3/1.95m ²						
	Filter membrane	Hydrophobic Polytetrafluoroethylene (PTFE)								
	Support layer		Polypropy	rlene (PP)						
Material	Housing/structural components		Polypropylene (PP)							
	Vent O-ring		Silicone							
	O-ring			Silicone/EPDM/FKM						
		Fo	orward:5.5bar(80psi)@25°C	Forward:5.5bar(80psi)@25°C						
	cimum operating erential pressure		1.7 bar(25psi)@80°C	1.7 bar(25psi)@80°C						
dille	erentiai pressure	R	everse:2.1bar(30psi)@25°C	Reverse:2.1bar(30psi)@25°C						
Maxir	mum inlet pressure		0.65Mpa							
]	Bubble point		≥1100mbar (100%, IPA	A bubble point@25°C)						
	Vater intrusion @2500mbar)	2.5 inches:≤0.2ml/min 5 inches:≤0.4ml/min 10 inches:≤0.8ml/min 20 inches:≤1.6ml/min								
Steril	lization resistance	Offline :126 °C /60 minutes, 3 times	Offline :126°C/30 minutes, 30 times (SIP not allowed)	Offline :126°C/60 minutes, 50 times SIP: Forward :142°C/30 minutes, 200 times Reverse :126°C/60 minutes, 25 times (not exceeding the maximum operating pressure difference of cartridge)						
	Cleanliness	Meet 21CFR21 0.3(b)(6) of "None fiber release"								
Bacterial endotoxin The endotoxin in the filtrate was determined according to the bacterial endotoxin test method specific the General Principles (1143) of the "2020 Chinese Pharmacopoeia". Bacterial endotoxin ≤ 0.25 EU										
I	Biosafety test	All materials of the filter element meet USP <88>, Class VI-70°C biosafety evaluation of plastic materials								
Ва	acteria retention	This product was tested for retention of Brevundimonas diminuta (ATCC 19146) by the ASTM F838 test method, and it's shown that the minimum retention rate was 10 ⁷ cfu/cm ²								
	Aerosol test	This product was tested for retention of Brevundimonas diminuta (ATCC 19146) aerosol referring to the requirements of PDA TR40, and it's shown that the minimum retention rate was 10 ⁷ cfu/cm ²								

Ordering information

Filgiant® Capsule filter





Filter Size:

D=20inch

E=30inch





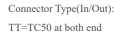


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Packing

Packing

Specification:

1= 1 per package

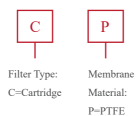
Specification:

1= 1 per package

Filgiant® Cartridge filter

Material:

P=PTFE



K

Filter Type:

K=Capsule



Filter Size:

A=2.5inch

B=5inch

C = 10inch

D = 20inch

E= 30inch

Membrane Pore Size: $02 = 0.22 \mu m$ $22 = 0.22 + 0.22 \mu m$



Sealing ring: S=Silicone E=EPDM F=FKM

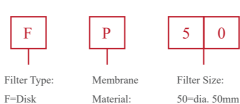






2=222 connector 1=internal insert connector

Filgiant® Disk filter

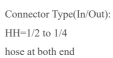


P=PTFE



Membrane Pore Size: $02 = 0.22 \mu m$ $22 = 0.22 + 0.22 \mu m$





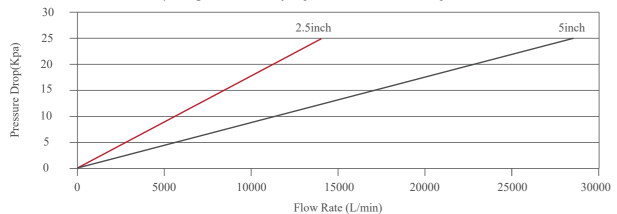


Packing Specification: 1= 1 per package

Filgiant® L series Hydrophobic Filter

Pressure drop and gas flowrate





Product parameters

I	Filter type	Capsule filter	Cartridge filter							
No	ominal size	1/2.5/5/10/20/30inch	2.5/5/10/20/30inch							
Effectiv	ve filtration area	$0.04/0.16/0.33/0.65/1.3/1.95m^2$	0.16/0.33/0.65/1.3/1.95m ²							
Meml	orane pore size	0.22μm/0.22+0.22μm/0.45μm/0.01μm								
	Filter membrane	Hydrophobic Polytetrafluoroethylene (PTFE)								
	Support layer	Polypropylene (PP)								
Material	Housing/structural components	Polypropyl	lene (PP)							
	Vent O-ring	Silicone								
	O-ring		Silicone/EPDM/FKM							
	num operating ential pressure	Forward:5.5bar(80psi)@25°C 1.7 bar(25psi)@80°C Reverse:2.1bar(30psi)@25°C	Forward:5.5bar(80psi)@25°C 1.7 bar(25psi)@80°C Reverse:2.1bar(30psi)@25°C							
Ві	abble point	0.22μm:≥1100mbar (100%IPA) ; 0.45μm:≥600mbar (70%IPA)								
ŗ	Water intrusion oure water C@2500mbar	2.5inch:≤0.2ml/min 5inch:≤0.4ml/min 10inch:≤0.8ml/min 20inch:≤1.6ml/min 30inch:≤2.4ml/min								
Steriliz	cation resistance	0.22μm SIP: Forward:142°C/30min, 100 times 0.22μm Offline:126°C/30min, 100 times 0.45μm Offline:135°C/30min, 5 times								
C	leanliness	Meet 21CFR21 0.3(b)(6)) of "None fiber release"							
Bacte	rial endotoxin	The endotoxin in the filtrate was determined according to the bacterial endotoxin test method specified in the General Principles (1143) of the "2020 Chinese Pharmacopoeia". Bacterial endotoxin ≤ 0.25 EU/ml								
Bi	osafety test	All materials of the filter element meet USP <88>, C	class VI-70°C biosafety evaluation of plastic materials							
Bact	eria retention	0.22μm: This product was tested for retention of Brevundimonas diminuta (ATCC 19146) by the ASTM F838 test method, and it's shown that the minimum retention rate was 10 ⁷ cfu/cm ²								
Insol	uble particles	Unless otherwise specified, the number of particl 1ml filter rinse solution shall not exceed 25 partic and more than 25 µm sha								

Ordering information

Filgiant® L Capsule filter





PolySaiTM Polypropylene (PP) Filter

Designed to remove particles from liquids and gases



Product features

- The PP filter pore size is calibrated with the real pore size, the wide range of pore sizes provides various choices for large and small particles retention
- PP is widely applied in liquid and gas filtration
- Wide pH tolerance from pH 1-14
- Strong particle-removing capacity, resistance to clogging
- No integrity test required

List of nominal pore size

- 0.1μm
- 5.0μm
- 0.3μm
- 6.0μm
- 0.5μm
- 10.0μm
- 0.6μm
- 20.0μm
- 0.8μm
- 50.0μm
- 1.0μm3.0μm
- 75.0μm100.0μm

Filter types

- Capsule filter
- Cartridge filter

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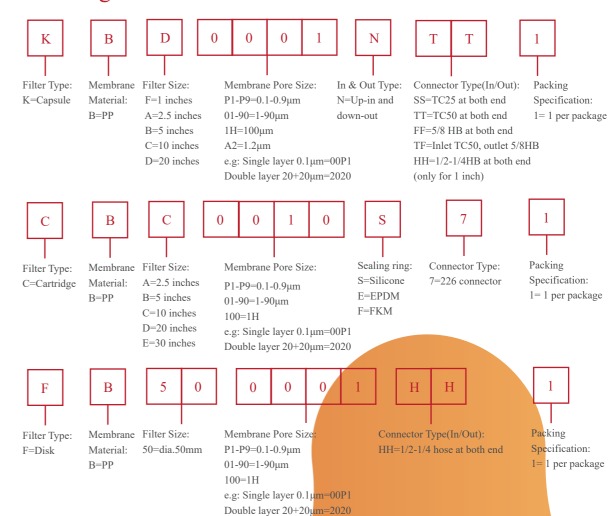
• Disk filter and customized filters



Product parameters

Filter type		Capsule filter				Cartridge filter					
Nominal size		1	2.5	5	10	20	2.5	5	10	20	30
Effective filtration	Single layer	0.038	0.16	0.33	0.65	1.3	0.16	0.33	0.65	1.3	1.98
area(m ²)	Double layer	0.035	0.14	0.28	0.55	1.1	0.14	0.28	0.55	1.1	1.65
Membrane pore size		0.1-100μm									
Filter membrane		Polypropylene (PP)									
Support layer		Polypropylene (PP)									
Housing/structural components		Polypropylene (PP)									
Vent O-ring		Silicone									
O-ring						Silicone/EPDM/FKM					
Maximum operating pressure		5.5bar(80psi)@25°C/1.7bar(25psi)@80°C									
Biosafety test		All materials of the filter element meet the biosafety evaluation of USP <88> class tests and Class VI-70°C plastic materials									

Ordering information



Peristaltic Pump

Peristaltic pump for 0.1m² TFF holder

Suitable for laboratory R&D and small batch production.

Uses brushless DC motor, suitable for pump head in series.

The upgrade model can realize the automatic allocation function of timing and quantity.

Product description

Dimensions: 285*207*180(MM)

Suitable hose specifications: 15# 24# 35# 36#

Maximum reference flow (ML/MIN): 6000(single pump head)

Basic version

Speed range (RPM): 60rpm-600rpm, reversible



Upgraded version

Speed range (RPM): 10rpm-600rpm, reversible





Fitting pump head

Torque wrench (10-50NM) Suitable for 0.1m² tangential flow filtration



Torque range (NM)	10-50
Driving head	3/8"
Length (MM)	426
Index value (N·m)	0.5

Hexagonal long sleeve



Φ(MM)	17
L(MM)	63.5

Peristaltic pump for 0.5m² TFF holder

With stainless steel housing, high protection level. Suitable for industrial production, using DC motor, suitable for pump head series. The upgrade model can realize the automatic allocation function of timing and quantity.

Product description

Dimensions: 380*326*214(MM)

Suitable hose specifications: 73# 82#

Maximum reference flow (ML/MIN):116000(single pump head)

Speed range (RPM): 60rpm-600rpm, reversible



 $Adapter\ pump\ head\ {\scriptstyle (Basic\ version)}$

Pump head housing material:



Adapter pump head (upgraded version)

Pump head housing material: stainless steel

Ordering information

TFTW01	10-50NM torque wrench		
TFTW05	20-100NM torque wrench		
TFP01E	0.1m ² peristaltic pump for fixture (economical)		
TFP01F	0.1m ² peristaltic pump for fixture (sub-assembly function)		
TFP05E	Peristaltic pump for 0.5m² fixture (economical)		
TFP05F	Peristaltic pump for 0.5m² fixture (sub-assembly function)		

Torque wrench (20-100NM) Suitable for tangential flow filtration of 0.5m² and above



Torque range (NM)	20-100
Driving head	1/2"
Length (MM)	473
Index value (N·m)	0.5

Hexagonal long sleeve (12.5 MM)



Φ(MM)	24
L(MM)	77

Validation Center

1.Product integrity test

In the production process, since most product/drug liquid contains materials or components which influences the surface interaction between filter membrane and the wetting liquid, the integrity test results can be depends on the cleanliness of those components. The product integrity test make it possible and validated to test the filter integrity based on the product liquid, since the the product liquid based integrity test and standard liquid based integrity test are data-associated.



2. Chemical compatibility test

Chemical compatibility test can be used to evaluate the chemical impact of the medium to be filtered on the filter under specific process conditions.



Membrane

Appearance, tensile strength, flux, thickness, SEM scan









Product	Test Project						
components	Appearance	Weight	Wire diameter	Tensile strengh	Flux	Integrity	
Menbrane	✓	✓	✓	✓	✓	✓	
Filter	✓				✓	✓	
Support components	✓						
Seal ring	✓	✓	✓				
TFF&NF	✓				✓	✓	

3.Extractables and leachtable test

Extractables refer to compounds that can be extracted from packaging or production component materials (plastics, elastomers and their coatings) under harsh experimental conditions (such as solvent, temperature and time) with solvents of different physical and chemical properties (solvents of different polarity or solutions similar to product liquid).

Leachable refers to compounds that migrate into a drug product from manufacturing parts, packaging parts, or delivery devices under normal manufacturing process, storage, and use conditions.

Basic testing process

- Carry out simulated extraction/simulated migration according to different component samples
- Speculate on potential extracts according to the results of simulated extraction
- Determine the risk substances according to the results of the extraction
- Carry out safety assessment according to risk substance content

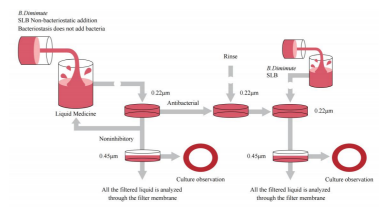
4. Bacterial retention test

Sterilizing filter

The sterilizing filter should be able to trap 1.0×107cfu of Btrevundimonas diminuta (ATCC 19146) by the ASTM F838 at a specific flow rate or pressure per square centimeter of effective filtration area.

Validation of bacterial retention process

To test the retention ability of sterilizing-grade filter with specific process fluid under specific filtration conditions, the bacterial retentation test is carried out on a laboratory scale under process conditions of scaling down.



Bacterial retention verification process

The purpose of the bacterial retention test is to simulate the worst conditions in the actual production filtration process, filter the product fluids containing a certain amount of challenging microorganisms or product substitute fluids, so as to confirm the microbial interception ability of the filter.

To test the retention ability of sterilizing-grade filter with specific process fluid under specific filtration conditions, the bacterial retentation test is carried out on a laboratory scale under process conditions of scaling down.

5. Validation service



center code



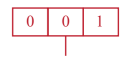
Laboratory Classification

ML=Microbiological Laboratory

VL=Virus Laboratory

EL=Chemical Laboratory

IL=Performance Laboratory



Test Project

002=Validation of bacterial retention process

004=Virus removal test

005=Extractables test and safety assessment

006=Leachable test and safety assessment

007=Chemical compatibility test

008=Product integrity test

009= Number of uses/lifetime verification

SaiflowTM Pharmaceutical-grade Stainless Steel **Filter Housing**

Adopt the latest sanitary design and the advanced polishing technology in the industry, and compatible with various types and brands of cartridge filter, so that it can be widely used in pharmaceutical, chemical, Bioengineering, food and beverage industries. The no-dead angle design inside the housing makes it more suitable for the use of high value-added feed liquid with very low residue.

Product features

- Manufacture in full compliance with cGMP standards and provide standard quality certificates
- Sanitary Design to meet regulatory requirements of the pharmaceutical industry
- Equipped with high point exhaust valve to ensure gas vent, prevent the formation of internal pressure or vacuum
- Equipped with low point exhaust valve to ensure liquid discharge, prevent the formation of residual and the condensed water discharge when the SIP
- Very low residue maximum recovery of product
- Provide customized design and manufacturing services

Product specification

Housing body	316L Stainless steel				
Gasket and sealing ring	Silicone/ EPDM/ FKM/ PTFE				
Clamp and feet	304 Stainless steel				
Connection type	Code 7 connection(2-226); Code 2 connection(2-222)				
In & Out Type	In line and T-line				
Sterilization form	SIP or autoclave				
Surface finish	Internal: Ra≤0.4μm; External: Ra≤0.6μm				
Housing size	1-cartridge housing: 2.5inch/ 5inch/ 10inch/ 20inch/ 30inc 2-cartridges housing: 2.5inch/ 5inch/ 10inch/ 20inch/ 30inc 3-cartridges housing: 2.5inch/ 5inch/ 10inch/ 20inch/ 30inc Other sizes can be customized				
Operating conditions	Pressure: -1 to 10 bar; Temperature: -10 to 150°C				

Ordering information

