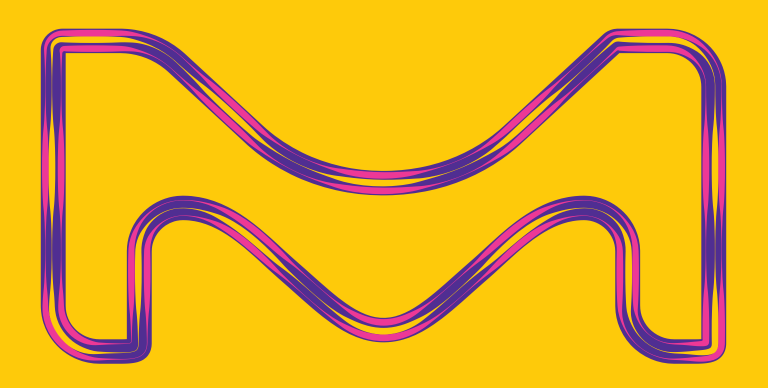


PARENTERAL PROCESS GUIDE

Formulation, Filtration, Filling





The life science business of Merck KGaA, Darmstadt, Germany operates as MilliporeSigma in the US and Canada.

SAFC®

Pharma & Biopharma Raw Material Solutions

Millipore®

Preparation, Separation, Filtration & Testing Products

BioReliance_®

Pharma & Biopharma Manufacturing & Testing Services



Reduce risk and increase efficiency with a fully assembled ready-to-use system:

- Let us help you design a process to meet your specific needs.
- What are the risks you are facing?
- What are your set-up and system issues to consider?



One-stop-shop to ensure component and process fit



Reduced risk of cross contamination and enhanced operator safety



Offering embedded in the Emprove® Program to facilitate regulatory compliance



Increase efficiency by reducing process and changeover time



Flexible manufacturing through:

- Reduced and flexible footprint allowing more free capacity
- Dedicated product and service offering



Quick link to technical and quality related solutions

Millipore®

The Millipore® portfolio of MilliporeSigma offers an ecosystem of industry-leading products and services, spanning preparation, separation, filtration and monitoring – all of which are deeply rooted in quality, reliability and timetested processes. Our proven products, regulatory and application expertise are a strong foundation you can rely on to consistently perform at the highest level.

Denotes Millipore® products

SAFC

The SAFC® portfolio of MilliporeSigma offers customized and ready-to-use raw material solutions, backed by deep regulatory expertise. Our high-quality products and services are supported by an experienced and responsive team of raw material and regulatory experts who are committed to understand your requirements and provide tailored solutions that meet your exact needs.

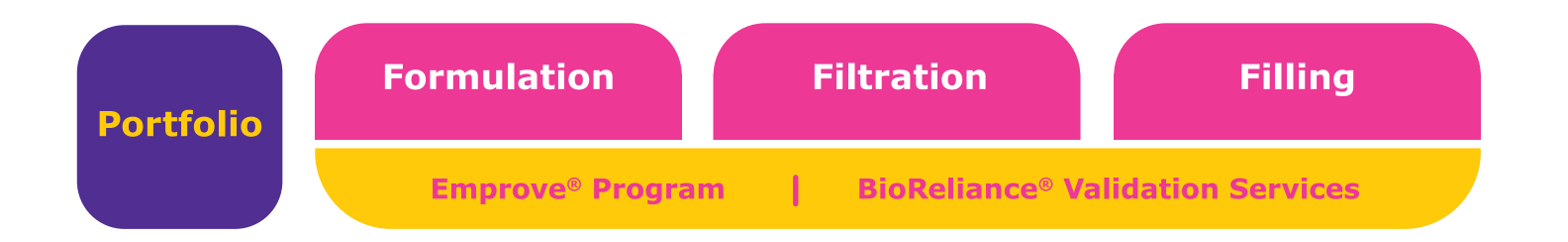
Denotes SAFC® products/services

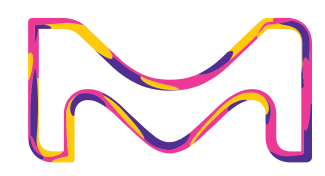
BioReliance®

The BioReliance® portfolio of MilliporeSigma encompasses biopharmaceutical characterization, safety testing and process development, as well as clinical and commercial biomanufacturing. Our experienced teams and operational expertise make us the partner who supports you all the way and always has your vital goal in mind.



Explore our portfolio >





Parenteral manufacturing process



Formulation



- Selecting high quality excipients
- Mixing of ingredients
- Monitoring critical parameters

Filtration

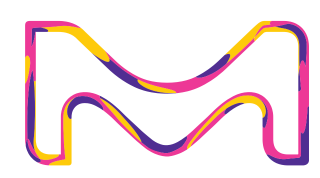


- Assure sterility of the drug product
- Contamination-free and closed sampling
- High recovery and low cost of goods

Filling



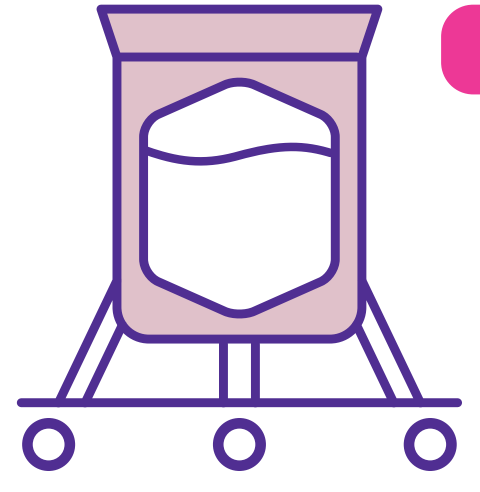
- Improve manufacturing flexibility
- Increase productivity
- Meet evolving manufacturing needs



Key steps for formulation

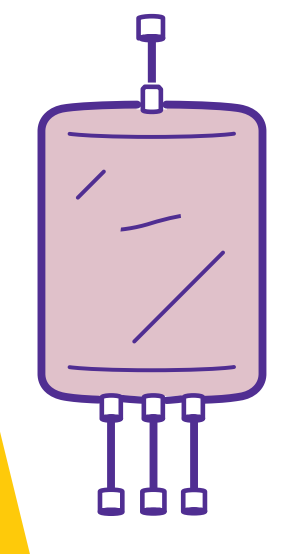
Select excipients

- Specifications and quality attributes designed for liquid formulations
- Reduce risk and simplify processes
- Reliable performance
- Ensure speed to market



Mixing

- Ensure mixing performance
- Gain flexibility through various working volumes and movability
- Choice for closed processing or open liners



Sterile sampling

- Improve operator safety
- Assure process sterility
- Sample representativity
- Ease of use







Considerations

Delivering Stateof the API

Ensuring Reliable Performance of the Final Drug Product

Market Readiness

Handling Powdered Raw Materials

Process Design

Solution

- Assess delivered state of API
- Aqueous
- Non-aqueous
- Co-solvent based
- Delivery system
- Sustained release
- Increased bioavailability, stability and solubility

- IPEC-GMP products assure consistent quality
- Reliable supply
- Biodegradable polymers for a sustained release profile
- Lipids with optimized characteristics for pharma applications





Considerations

Delivering State of the API

Ensuring Reliable Performance of the Final Drug Product

Market Readiness

Handling Powdered Raw Materials

Process Design

Solution

Choose

- Buffers
- Mineral salts & tonicity
- Preservatives
- Solvents
- Surfactants
- Antioxidants
- Stabilizers

- Comprehensive product portfolio
- Broad range of IPEC-GMP products
- Support during development, scale-up and commercial production
- Batch-to-batch consistency





Considerations

Delivering State of the API

Ensuring Reliable Performance of the Final Drug Product

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Handling Powdered Raw Materials

Process Design

- i i occas Designi
- Meeting quality requirements
- Ensuring speed to market
- Documentation for regulatory approval

- Solution
- Reduce risk and simplify processes through <u>the Emprove</u>® <u>Program</u>
- Low Bioburden, Endotoxins and Reducing Sugars
- IPEC-GMP Manufacturing
- Multicompendial





Considerations

Delivering State of the API

Ensuring Reliable Performance of the Final Drug Product

Market Readiness

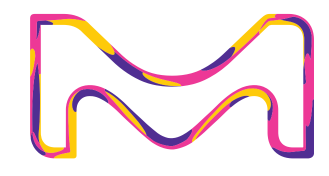
Handling Powdered Raw Materials

Process Design

Solution

- Minimized product contamination risk
- Increased personnel and facility safety

- Paper free packaging to control the particulate matter
- Packaging system to minimizes
 product caking
- Granulated excipients for better processing and handling



How does the Emprove® Program help you

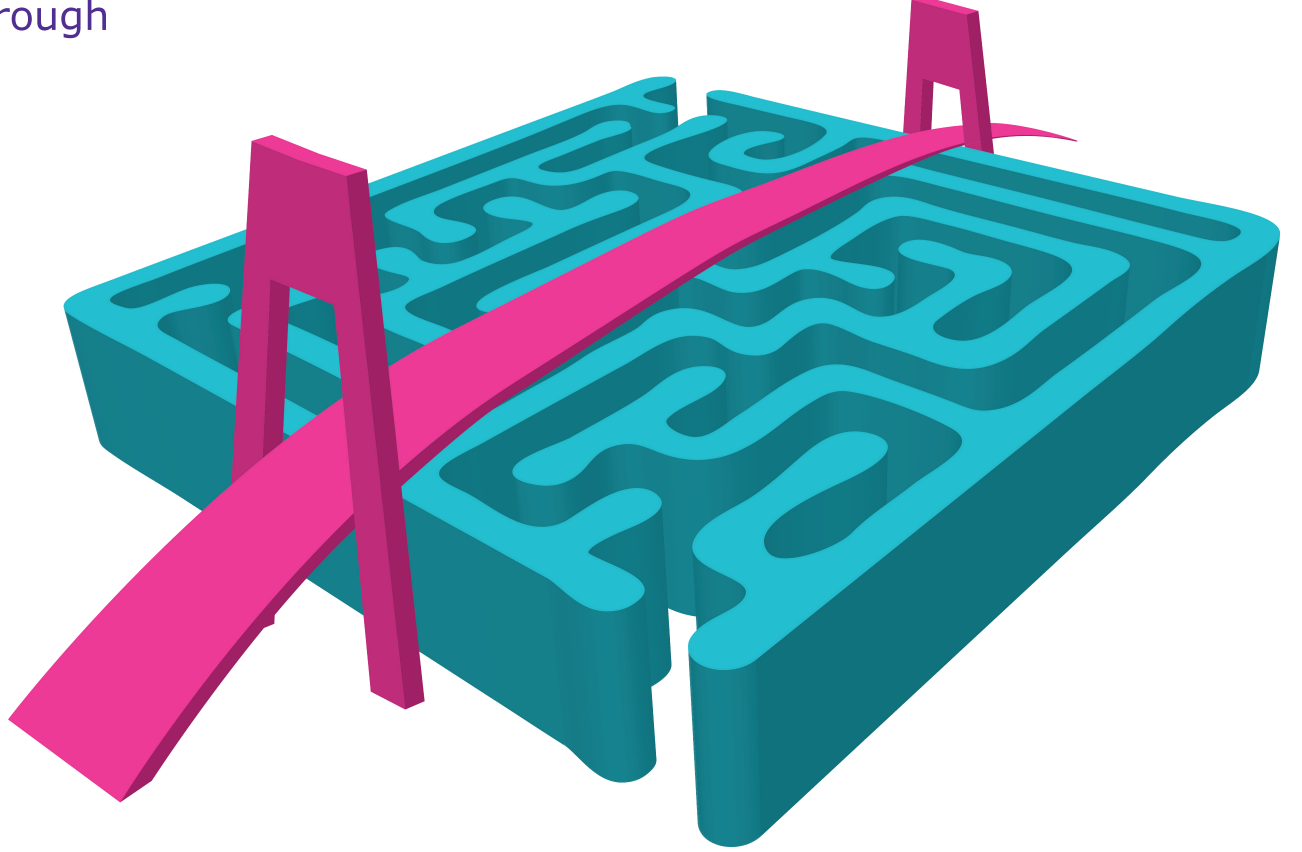
Qualification, risk assessment, and process optimization

The Emprove® Program contains over 400 raw and starting materials as well as 20 filter and single-use product families. Each product portfolio is supported with Emprove® Dossiers which provide comprehensive and up-to-date documentation to help you navigate regulatory challenges, manage risks, and improve your manufacturing processes.

Emprove® Dossiers are available exclusively through the Emprove® Suite subscription.

Emprove® Dossiers

- Material Qualification
- Quality Management
- Operational Excellence





Exemplary specification parameters of two of our stabilizers

Reducing sugars are normally incompatible with the primary amine. That leads to discoloration of the solution and reduction of the potency.

Endotoxins are the most significant pyrogens in parental drugs and the cause of strong immune responses.

To control the reducing sugars and the endotoxin levels in the raw materials, we offer lower limits than Pharmacopeia specifications.

Sucrose	
low in endotoxing	S
≤ 0.3 I.U./g	

	Endotoxins (I.U./g)		Reducing Sugars (%)	
	SAFC®	Ph Eur, USP	MilliporeSigma	Ph Eur, USP
Sorbitol	≤ 1	< 2.5	≤ 0.11	≤ 0.2
Mannitol	≤ 1	< 2.5	≤ 0.05	≤ 0.1

Packaging

Handling

DRYPOUR™ – offers triple protection of the material: against moisture and contamination from outside and absorption of moisture present in the material

- Polyethylene (PE) drum with tamperevident seal
- Polyethylene inliner with integrated desiccant bags
- Tyvek[®] inliner



Dispense

Increase product and operator safety with bulk powder transfer packaging

- Direct dispense solution
- Customizable quantity
- Ergonomic design



Packaging

Most raw materials are available in paper-free packaging including

- Pails
- Drums
- Pre-packs
- Double PE bags
 - Specially-designed for cleanroom manufacturing
 - Removal of the outer bag outside of the clean room
 - Labelled inner bag to minimize handling risks



- **Buffers**, to ensure best pH for physiological tolerance, solubility and stability
- Mineral salts & tonicity adjustment, to ensure the osmolarity for physiological tolerance
- **Preservatives**, to ensure microbial purity and sterility by inhibiting microbial growth
- **Solvents & surfactants**, to enhance solubility and stability of the API
- Antioxidants, to ensure the stability of oxidation prone APIs
- Stabilizers, to stabilize the finished drug product







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Choosing the right mixers for homogenous and effective formulations

Considerations

Application Requirements

Operator Friendly Solutions

Facility Needs

Validation

Process Design

Solution

- Sensitive formulation
 - Small molecule: cytotoxic and hormonal drugs
 - Peptides and protein solutions
- Low to high concentrations
- Difficult to mix solutions
 - Polymers
 - Oil-based formulation

- Low shear gentle mixing (liquid-liquid mixing)
- High mixing strength creating vortex (solid-liquid mixing)
- Optimization, technical and process support by team available
- Live demonstration in the M Lab™





Choosing the right mixers for homogenous and effective formulations

Considerations

Application Requirements

Operator Friendly Solutions

Facility Needs

Validation

Process Design

Solution

- Plant specific requirements
- Convenient powder delivery
- In-process measurement options
- Representative sampling
- Direct filtration

- In process measurements parameters:
 pH, weight, temperature, conductivity
- Powder delivery system
- Closed, sterile, zero-dead-leg sampling
 system





Choosing the right mixers for homogenous and effective formulations

Considerations

Application Requirements

Operator Friendly Solutions

Facility Needs

Validation

Process Design

Solution

The conceptual design

- Process economics
- Flexibility for product change-over
- Purpose of facility
- Ease of process validation

Benefits of single use mixing systems:

- Decreased risk of contamination
- Reduced downtime and costs for cleaning validation
- Reduced capital
- Faster turn-around
- Flexibility to change scale or process
- Closed processing for critical applications or open liners for lower value applications

Pharma & Biopharma Manufacturing & Testing Services



Choosing the right mixers for homogenous and effective formulations

Considerations

Application Requirements

Operator Friendly Solutions

Facility Needs

Validation

Process Design

Solution

Equipment coming in contact with the product assembly must not be

- Additive
- Reactive
- Absorptive

- Validation team experts
 - To help you choose and conduct the appropriate validation services for your mixing process
 - To accelerate and simplify your path to market
- Documentation support to meet your compliance needs



How is the Mobius® Power MIX different than the Mobius® MIX?

Mobius® MIX

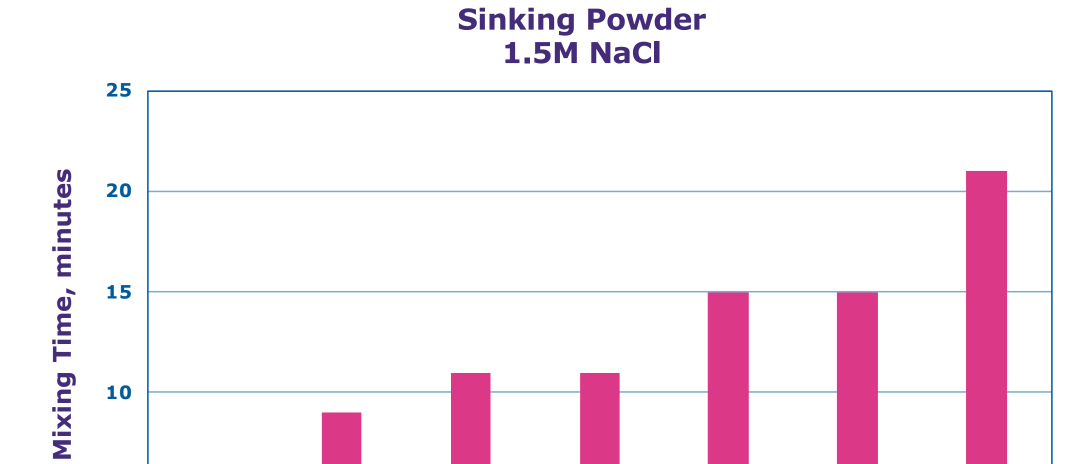
Gentle mixing of drug products throughout entire process.

Mobius® MIX

Levitating, magnetically driven impeller

- Low shear impeller design
- No particle shedding
- Mixing from 10 L 1000 L

Mobius® Power MIX Forms a vortex for powerful mixing of difficult-to-mix solutions.



500

1000

Mobius® Power MIX Size

2000

2500

3000

100

200

Scalability of mixing

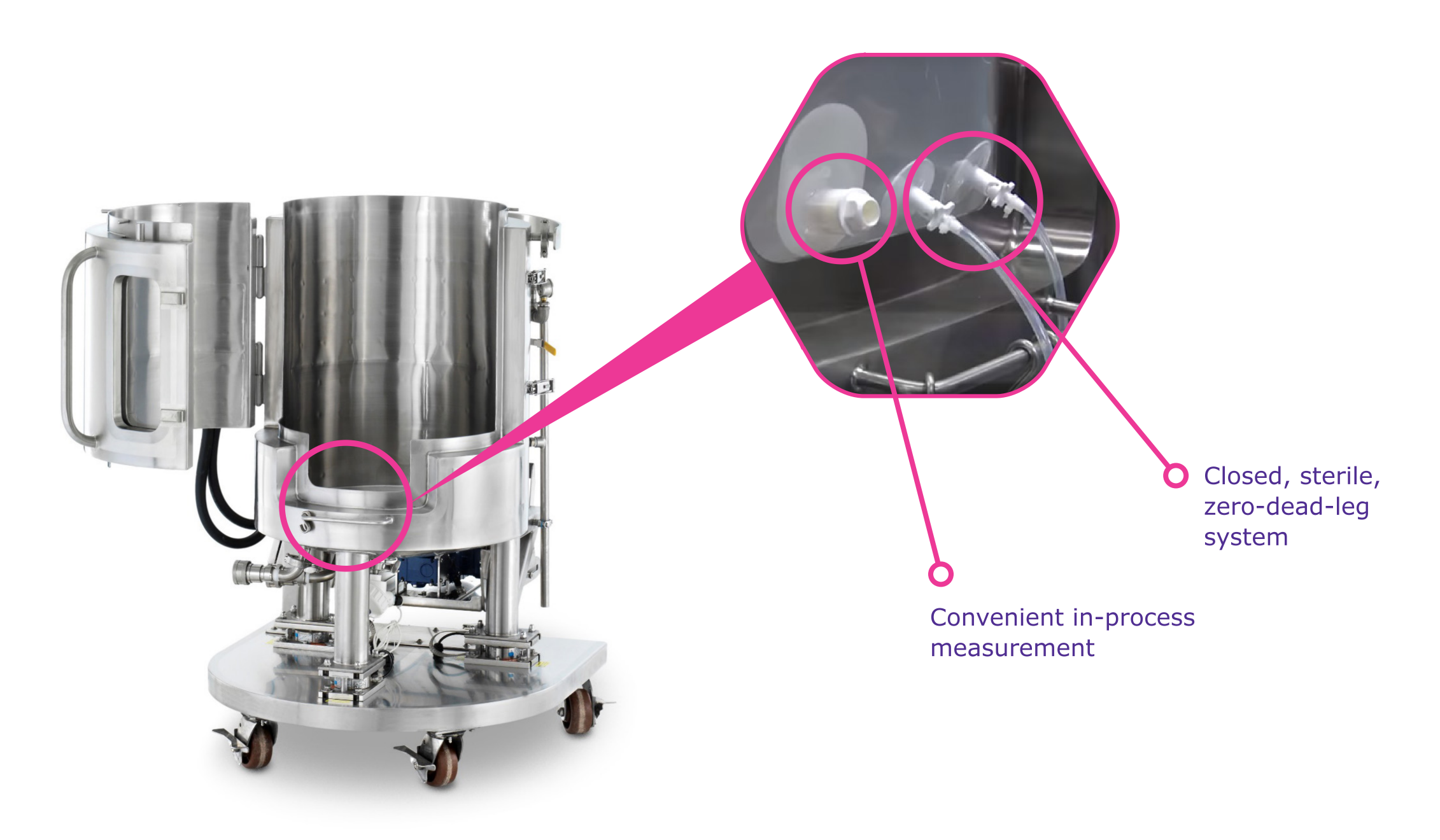
Mobius® Power MIX

Magnetically driven impeller (NovAseptic technology)

- More efficient powder/liquid mixing
- Viscous solutions
- Mixing capabilities from 100 L 3000 L



Inline measurement and sampling



BioReliance® Validation Services

Validation services for mixing bags

Emprove® Program

Chemical compatibility

Assess the chemical compatibility based on key characteristics, after prolonged exposure with the drug product. Provide evidence that the process fluids and conditions do not adversely impact the structure of the mixing bag.

Patient safety

Assess the potential impact of the substances that have been detected, identified and quantified on patient safety. This assessment is done by a toxicologist.

Extractables

Identify and quantify the extractables which may be extracted out from the mixing bag by employing the Model Solvent Stream Approach and worst case test conditions. Analytical methods used are:

NVR: Non-Volatile Residue

TOC: Total Organic Carbon

FTIR: Fourier Transform Infrared

Spectroscopy

RP-HPLC: Reverse-Phase High-Performance Liquid Chromatography

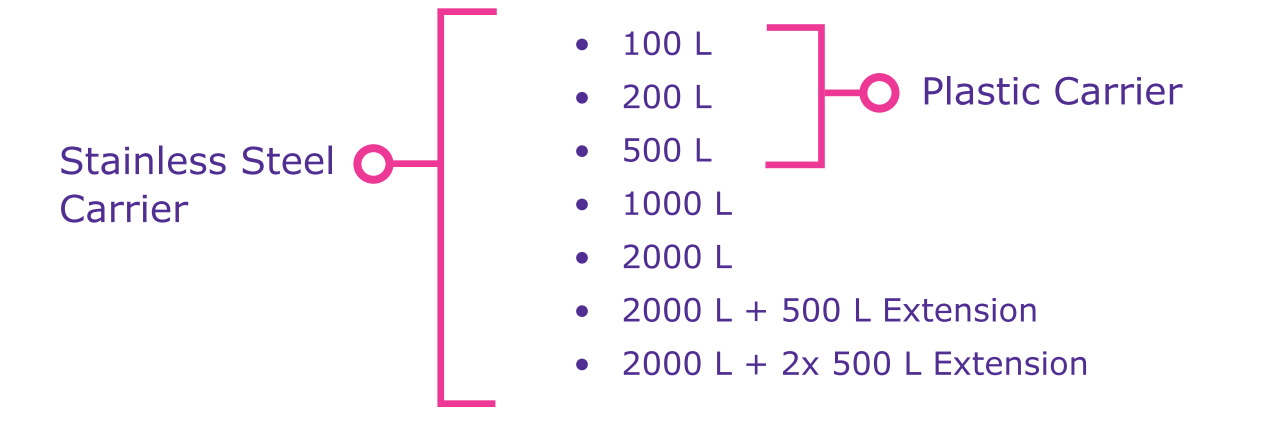
GC-MS: Gas Chromatography/ Mass Spectrometry

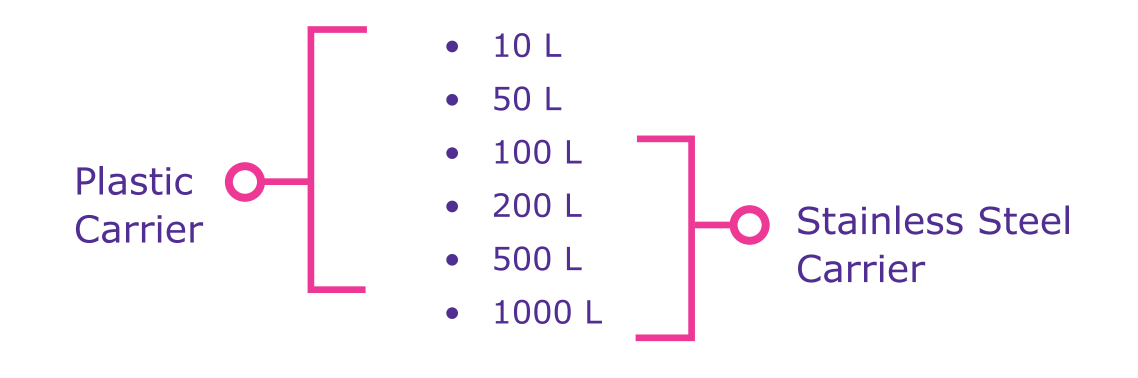


BioReliance® Validation Services



Mobius® Power Mix



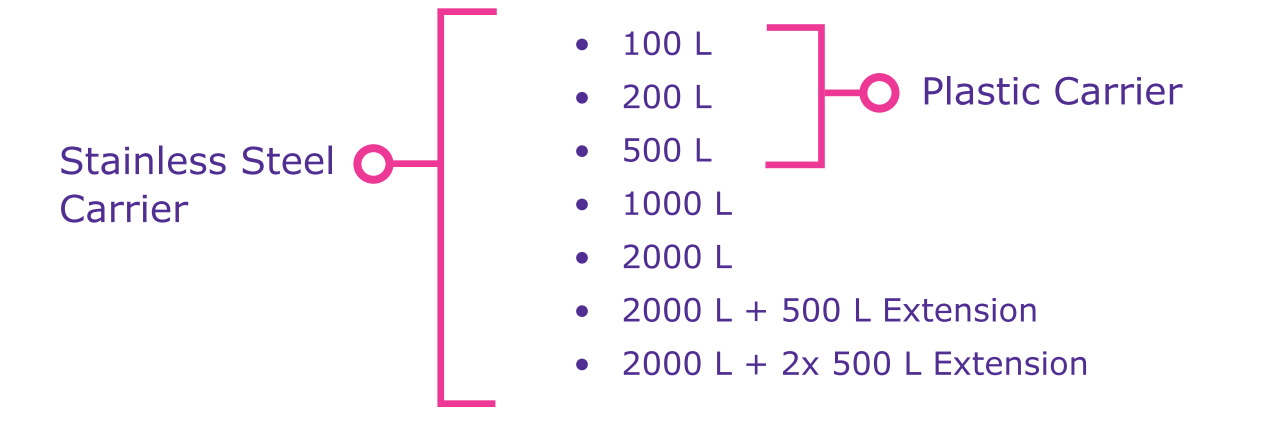


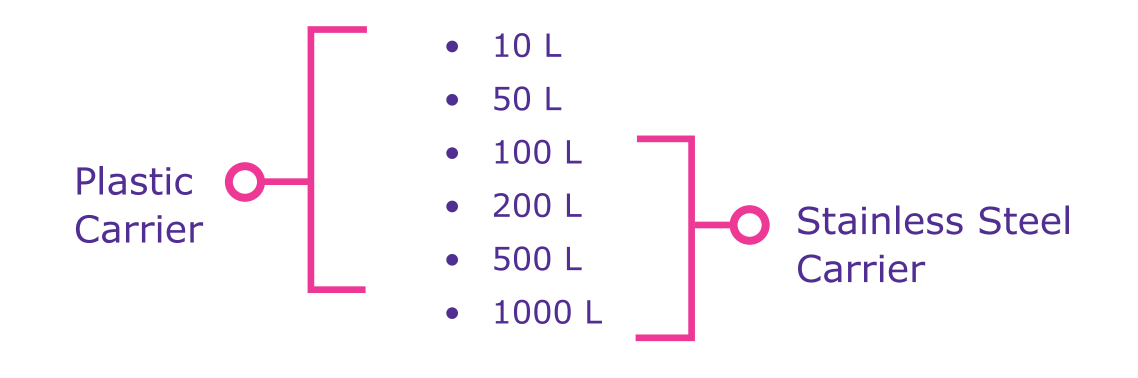


BioReliance® Validation Services



Mobius® Power Mix



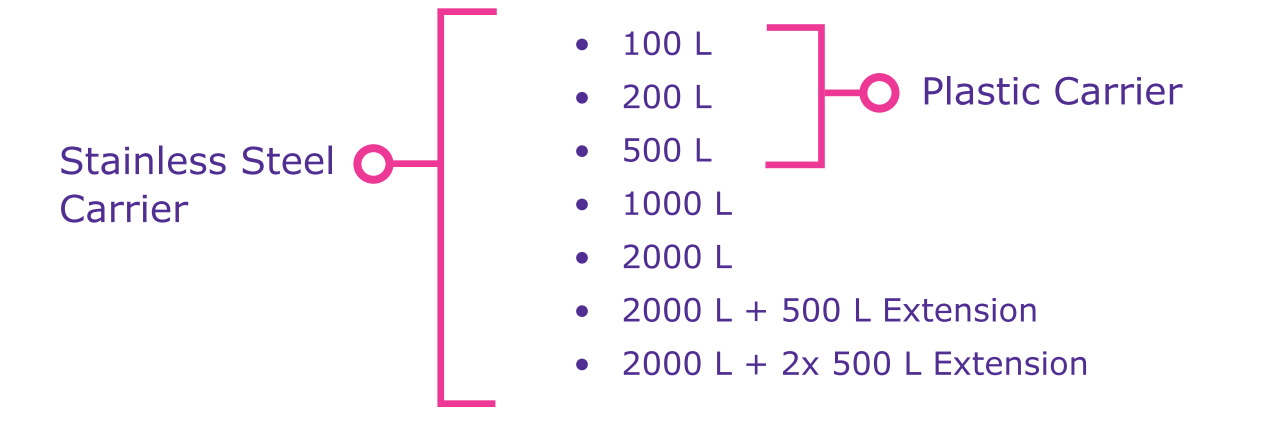


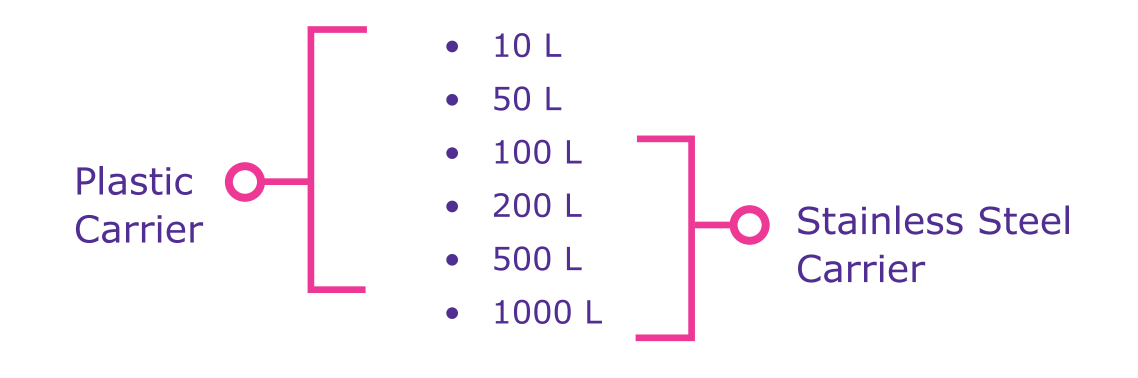


BioReliance® Validation Services



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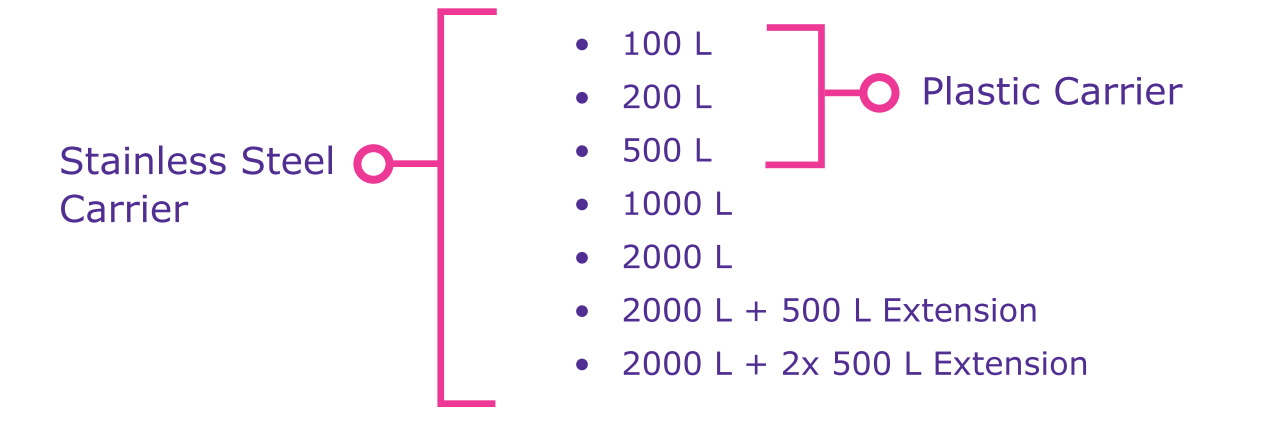


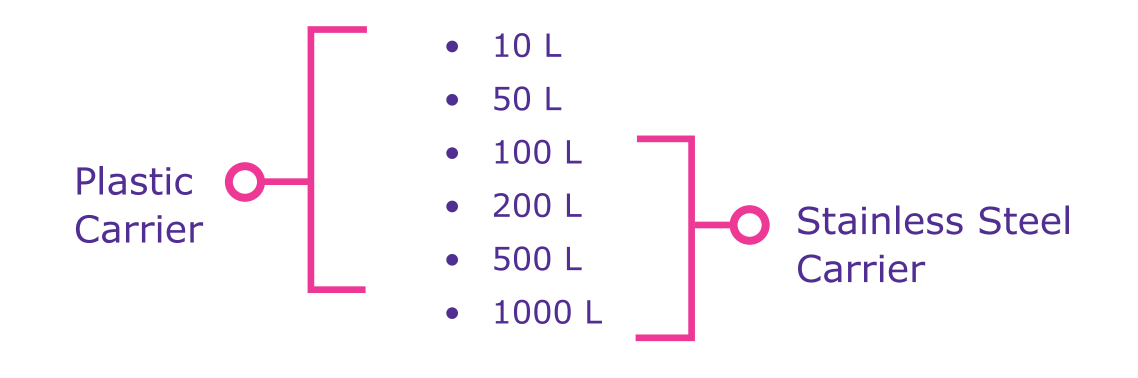


BioReliance® Validation Services



Mobius® Power Mix











Considerations

Ease of Use Facility Needs Meeting Regulation Validation Requirements **Process Design** Solution Solutions for all process steps Flexibility and speed Safety of operators Wide range of sampling containers, Secure disconnection of sample available pre-sterilized and pre-assembled Preconfigured or configure on site No operator training needed





Choosing the right sampling systems to monitor critical parameters across each step

Considerations

Ease of Use

Facility Needs

Meeting Regulation Requirements

Validation

Process Design

Solution

- No need for additional utilities (pressure, steam, ...)
- Option of sampling from SU and SS
- Limited investment
 - Maintenance
 - Spare parts

- Utilities: only need of crimping tool to disconnect taken sample
- Sampling from single-use and stainless steel tank possible
- Reduced cleaning needs





Choosing the right sampling systems to monitor critical parameters across each step

Considerations

Process Design

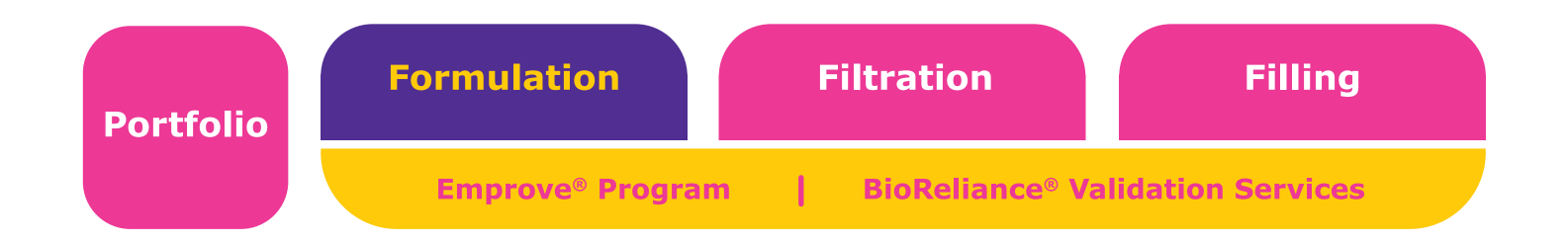
Process Design

Solution

Process integrity
Representative samples
Process risk mitigation
Accelerate speed to market

Meeting Regulation

Process integrity
No dead leg, flush or sample dilution
Closed and validated sampling system





Pharma & Biopharma Manufacturing & Testing Services





Choosing the right sampling systems to monitor critical parameters across each step

Considerations

Ease of Use

Facility Needs

Meeting Regulation Requirements

Validation

Process Design

Solution

Equipment coming in contact with the product must not be

- Additive
- Reactive
- Absorptive

- Validation team experts
 - to help you choose and conduct the appropriate validation services for your sampling system
 - to accelerate and simplify your path to market
- Documentation support to meet your compliance needs

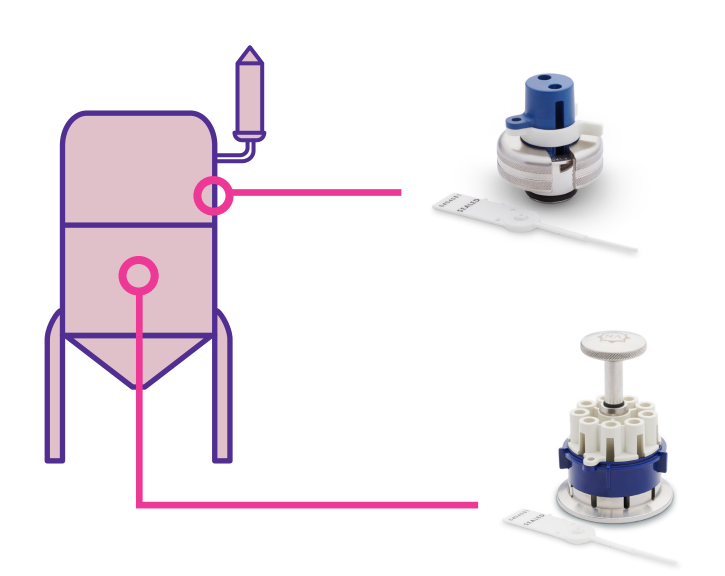


Emprove® Program | BioRel

BioReliance® Validation Services

Process step options

Connector for stainless steel tank



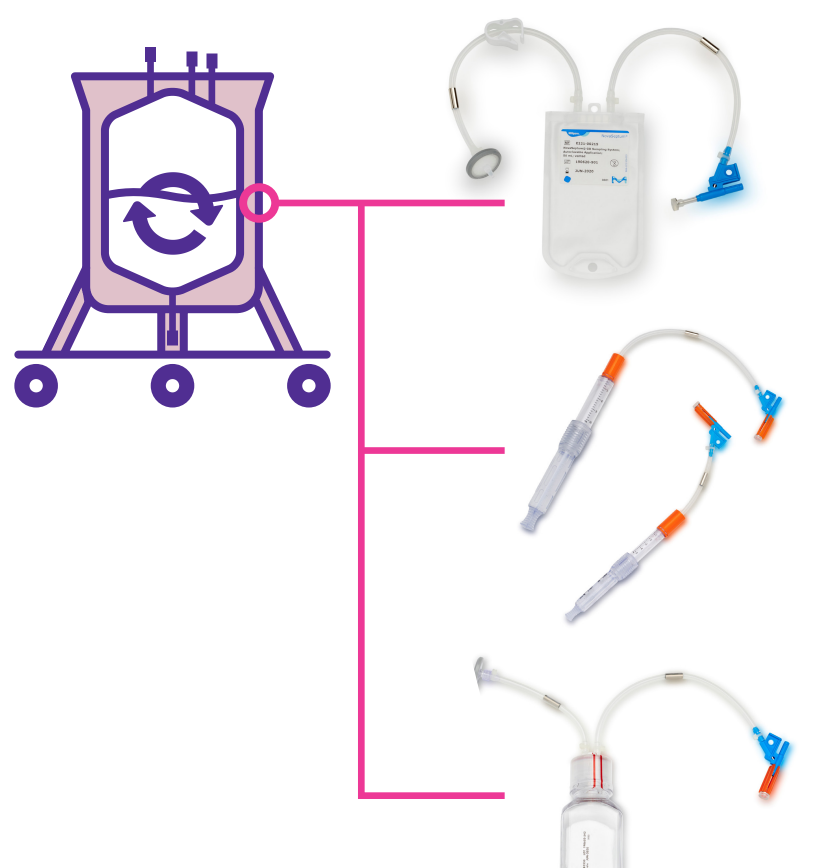
Ingold connectors

- Available in any length
- 2 independant points of access

TC connectors

- 1/2" to 2" size with up to 10 independent access points
- Also available in singleuse format, pre-assembled with the containers of your choice

Sampling containers



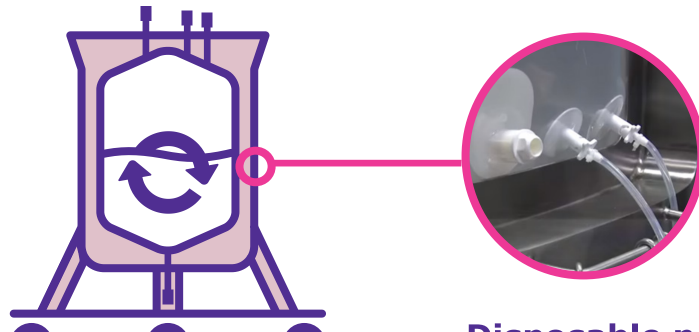
Autoclave containers

- Bags and syringes ideal for small portable tanks that are fully autoclaved
- Vented to withstand the most aggressive autoclave cycles

Sterile syringes

- Contamination free with patented design
- Ideal to save and store high value product

Connector for single-use tank



Disposable needle-free valve

- Ideal for sampling from disposable bags
- Welded any place needed

Bottles

- Bio-neutral plastic bottles ideal for all tests
- Offered in multiple sizes

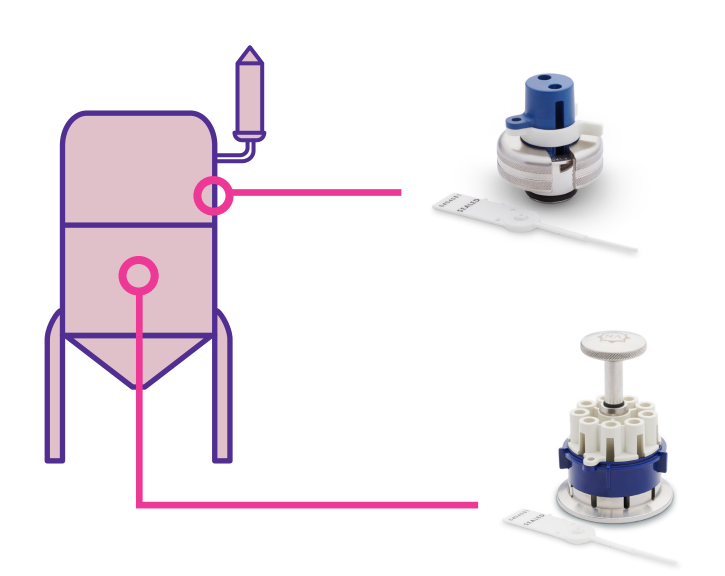


Emprove® Program | BioRel

BioReliance® Validation Services

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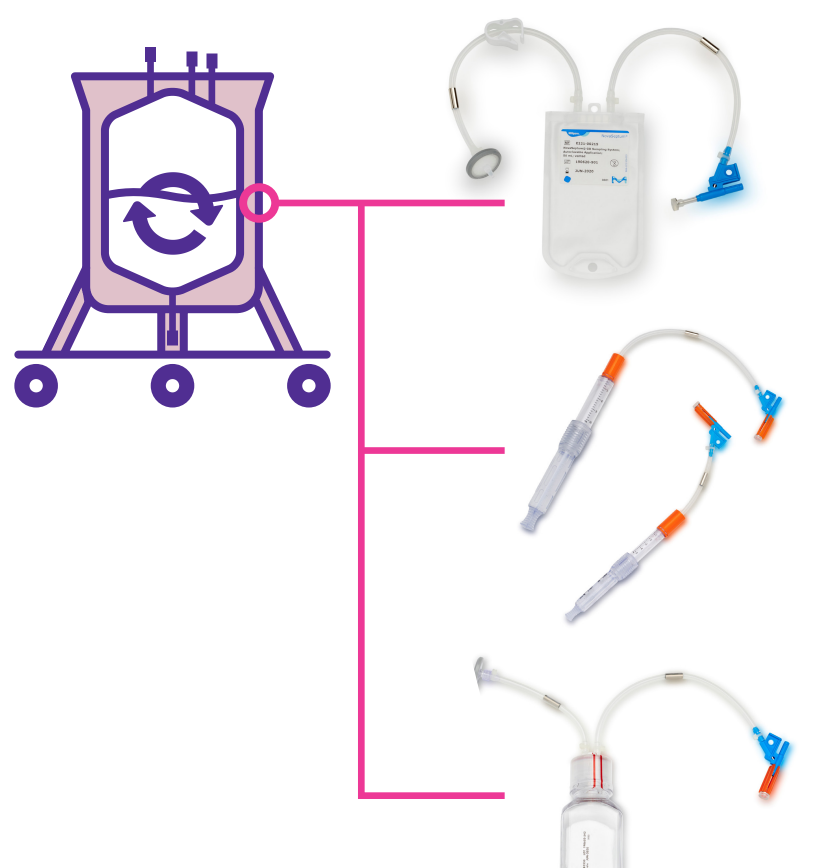
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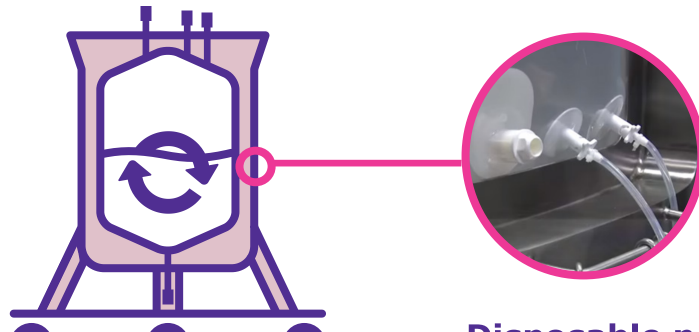
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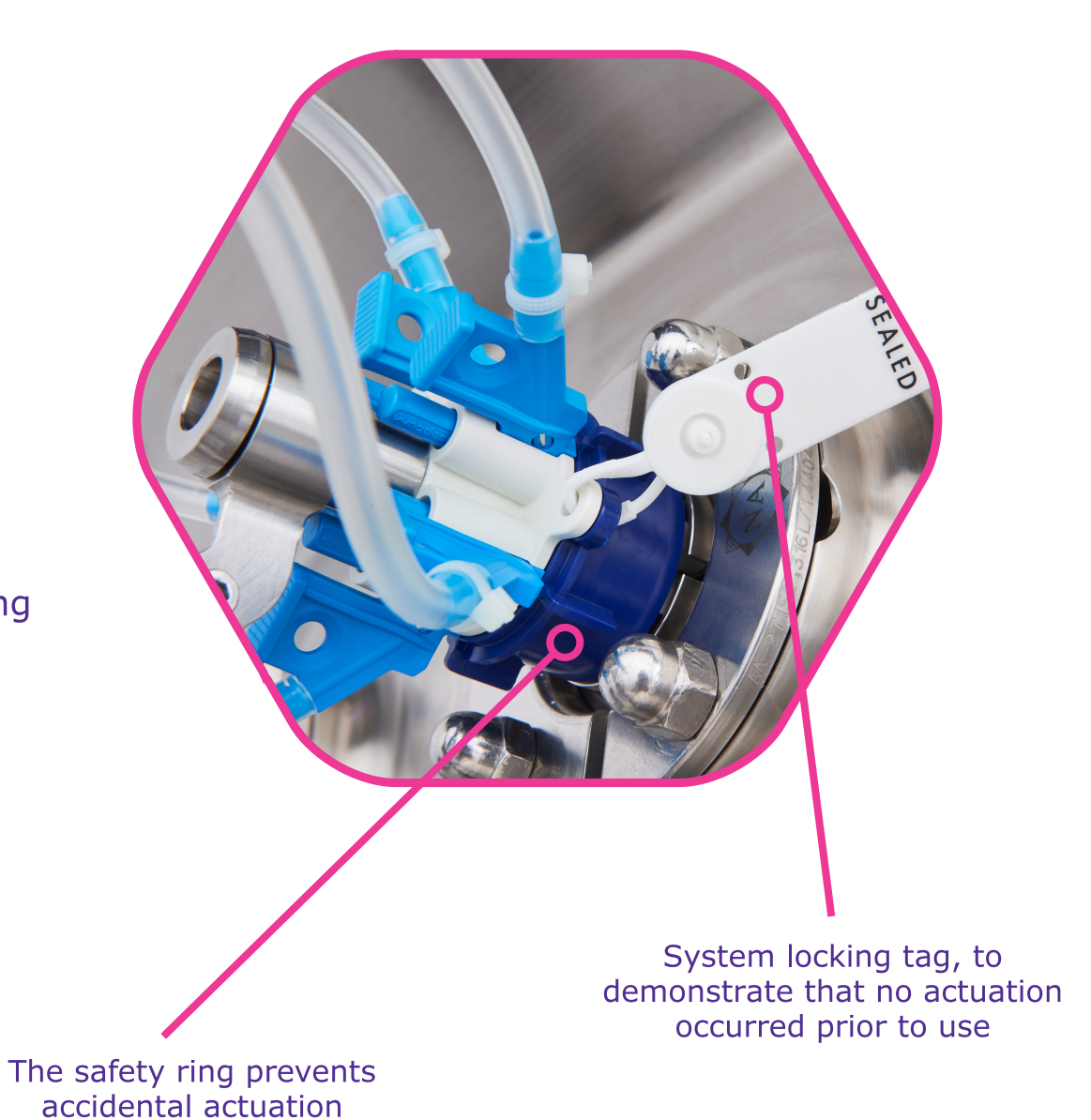
during processing

Emprove® Program | BioRe

BioReliance® Validation Services

Maximum security and reliability

- Eliminating high heat/pressure and glass bottle safety risks
- No steam condensate to dilute sample, avoid requirement for flushing
- "Neutral" containers for accurate testing and storage
- Simple procedure eliminates complexity of training, reducing risk of operator bias
- Eliminating time of steam sterilization and cooling between samples



Validation services for sampling system

Chemical compatibility

Assess the chemical compatibility based on key characteristics, after prolonged exposure with the drug product. Provide evidence that the process fluids and conditions do not adversely impact the structure of the Sampling System

Functionality test

Verify that the sampling element is within the qualified acceptance criteria after product contact & process conditions simulation.

Non permeation test for bag

Demonstrate that the bag is not permeable to disinfectant/ decontamination agent



NovaSeptum® GO sterile sampling holders and connectors

BioReliance® Validation Services

Multi-Use holders for stainless steel

Emprove® Program

Portfolio

 A fast and easy connection of the sampling solution to the manufacturing process available.



Ingold® Holder



TC Holder

Single-Use connectors for single-use assemblies

- Reduce labor associated with cleaning or handling.
- with NovaSeptum[®] single-use connectors.
- The needle-free sampling valve brings capabilities for NovaSeptum® sampling integration into your larger single-use bag assemblies.



Single-use holder for stainless steel

 Available pre-loaded with high purity or autoclavable bags, bottles or AV syringe





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Single-use holder for stainless steel

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The NovaSeptum® GO high purity bag single and multi-sampling systems are available from 50 mL to 1000 mL.



The NovaSeptum® GO autoclavable bag single sampling system is available in 50 mL, 100 mL, 250 mL and 1000 mL.



The NovaSeptum® GO bottle single and multisampling systems are available in 60 mL to 500 mL.



The NovaSeptum® GO transfer unit is available in different tubing materials (Thermoplastic Silicone, and C-Flex®).



The NovaSeptum® GO AV single and multisampling units are available from 5 mL to 20 mL syringe sizes.





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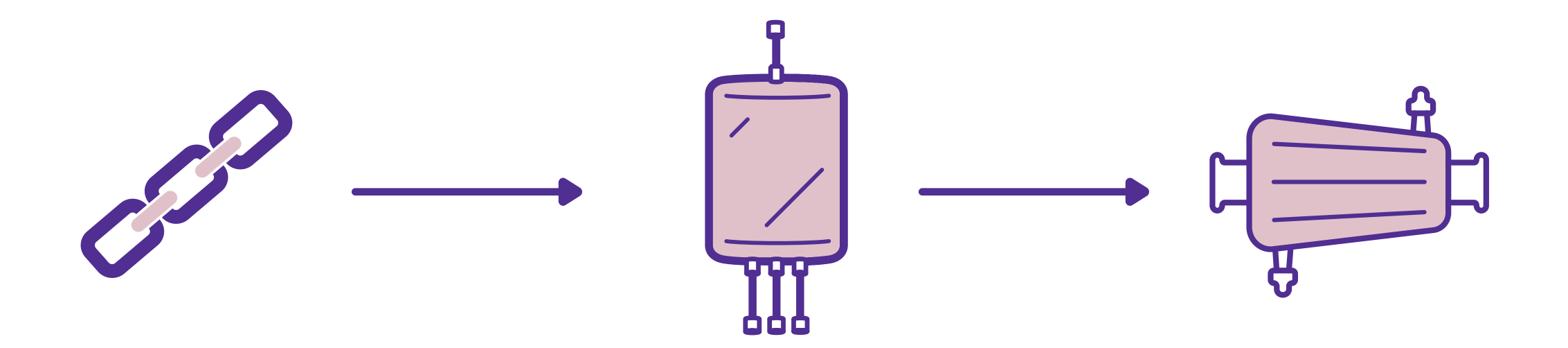
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Key steps for filtration



Sterile connection •

- Assure the fluid path is not compromised between process steps
- Transition between multi-use and single-use systems

Sterile sampling

- Improve operator safety
- Maintain process sterility
- Assure representative sampling
- Ease of use

Sterile filtration

- Filters designed for critical sterile filtration steps
- Expertise in single-use filtration system design



Chosing the right sterile connectors to maintain an aseptic fluid path

Considerations

Sterility Assurance Application Validation Handling **Process Design** Solution Quick and easy connectivity Robust design delivers consistent and reliable performance Maintain sterile flow-path Avoid operator mistakes Operator handling training



Chosing the right sterile connectors to maintain an aseptic fluid path

Considerations

Handling **Validation Sterility Assurance Application Process Design** Solution Completely closed fluid path 100% air-integrity tested in Sterile connection could be made in both manufacturing > non-classified and classified area Aerosolized microbial challenge test – Eliminate contamination risk associated worst case simulation for validation > with aseptic connections



Chosing the right sterile connectors to maintain an aseptic fluid path

Considerations

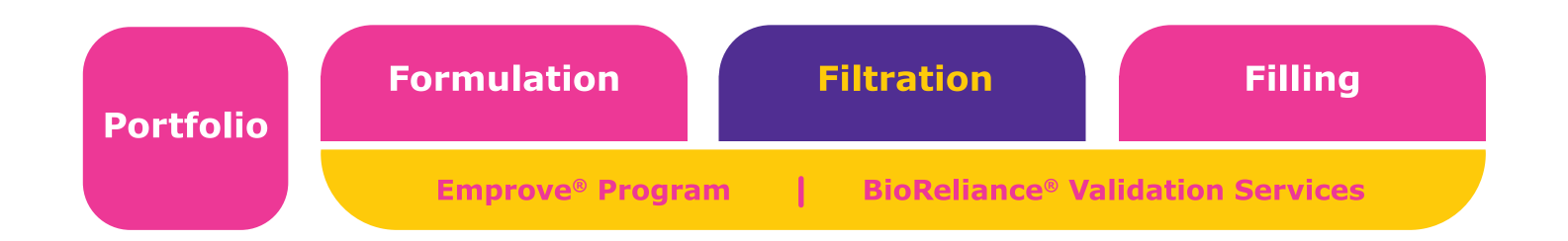
Handling Sterility Assurance Application

Validation

Process Design

- Sterile fluid transfer
- Connect sterilized fluid paths, integrate steamable hard-piped connections to disposable fluid paths
- Single and multiple actuations

- Solution
- Sterile-to-sterile connection
- Steamable hard-piped connection to disposable fluid path
- Multiple connections/disconnections







Chosing the right sterile connectors to maintain an aseptic fluid path

Considerations

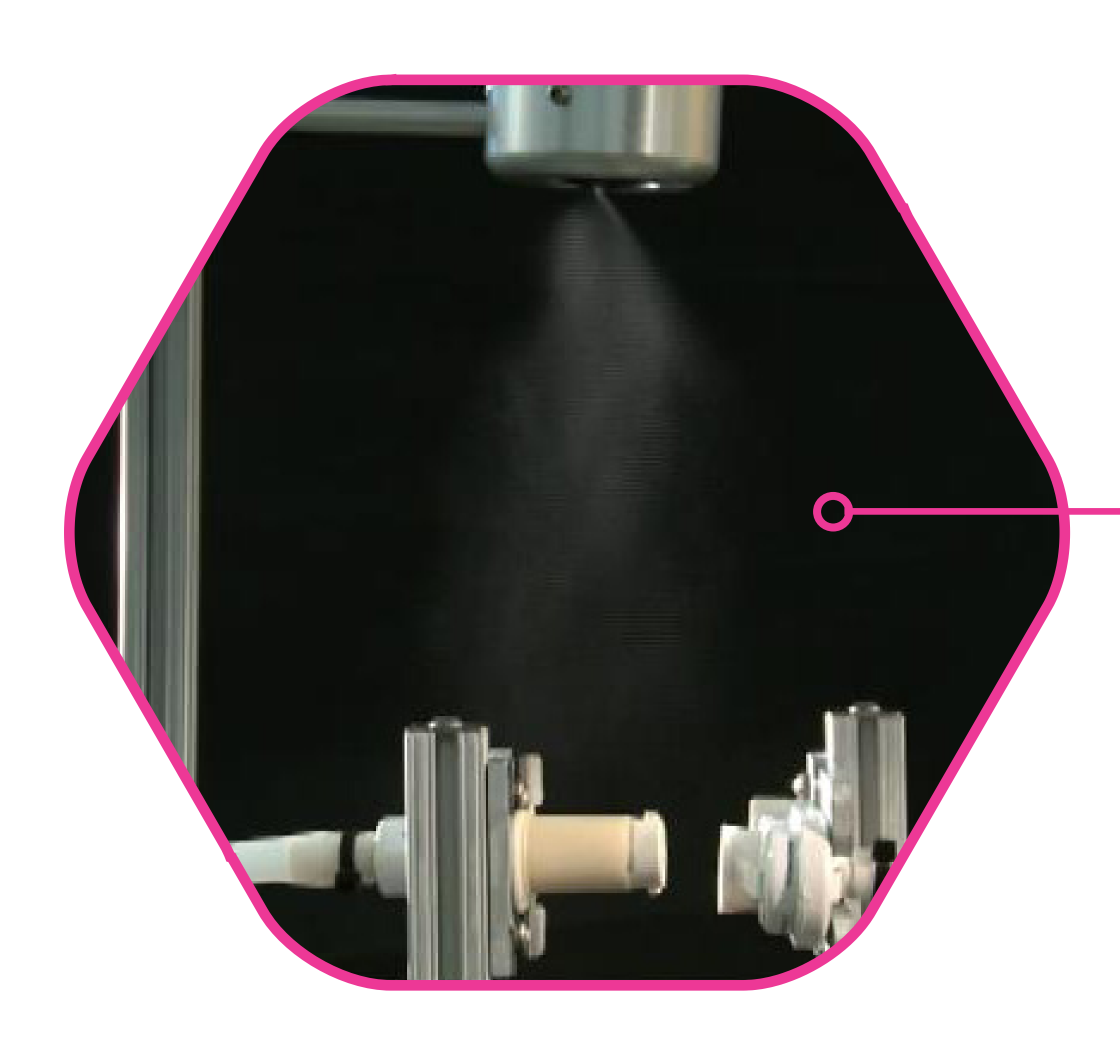
Handling Sterility Assurance Application Validation

Process Design Solution

Equipment coming in contact with the product must not be

- additive
- reactive
- absorptive

- Validation team experts
 - to help you choose and conduct the appropriate validation services for your sterile connectors
 - to accelerate and simplify your path to market
- Documentation support to meet your compliance needs



Brevundimonas diminuta, >4 x 10⁶ cfu per connector set

Sterility assurance during actuation

Two tests were conducted to challenge the ability of Lynx® S2S connectors to prevent ingress of microorganisms in a non-classified area.

Tests were performed to confirm that the Lynx® S2S connectors showed no ingress of the challenge organism, *B. diminuta*.

Bacterial aerosol

The connector was connected and actuated in the presence of a bacterial aerosol at a minimum concentration of 1×10^6 cfu/connector.

Direct bacterial soiling

B. diminuta was applied to the mating surfaces of the male and female Lynx® S2S connector lumen plugs, followed after a setting time, by connection and actuation.

All Lynx® S2S connectors met the acceptance criteria for the Bacterial Aerosol Test and the Direct Bacterial Soiling test, assuring a sterile connection can be made in non-classified areas.

Validation services for sterile connectors

Chemical compatibility

Assess the chemical compatibility based on key characteristics, after prolonged exposure with the drug product. Provide evidence that the process fluids and conditions do not adversely impact the structure of the connectors.

Assess the potential impact of the substances that have been detected, identified and quantified on patient safety. This assessment is done by a toxicologist.

Patient safety

Extractables

Identify and quantify the extractables which may be extracted out from the connectors by employing the Model Solvent Stream Approach and worst case test conditions. Analytical methods used are NVR, TOC, FTIR, RP-HPLC and GC-MS (when applicable).

> Click here for more information on the **BioReliance® Validation Services**

BioReliance® Validation Services



Lynx® product family







Lynx® S2S

Sterile-to-Sterile Connector

A single actuation, disposable device for connecting sterilized disposable flow paths

Lynx® ST Steam-To Connector

Designed to connect steamable hard-piped processing systems to sterilized disposable flow paths

Lynx® CDR

Connect, Disconnect, Reconnect

BioReliance® Validation Services



Lynx® product family







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Choosing the right filters to sterilize your product

Considerations **Line Flushing Validation** Filter design **Filtration System Integrity Testing** Setup **Process Design** Solution Streamlined single-use assembly Maximizing yield, minimize loss Ease of use Truly closed systems Membrane selection (sterility assurance) Reliable sterile filtration of small volume and high value solutions > level, low adsorption, flow rates, capacity, chemical compatibility) Maintain sterility of flow path







Choosing the right filters to sterilize your product

Considerations

Filter design

Filtration System Setup

Line Flushing Integrity Testing

Validation

Process Design

Solution

- Single Dual Redundant Filtration
- Line orientation (vertical, sloped, or horizontal)
- Sterile Filter placement (inside or outside of the isolator)
- PUPSIT implementation (Pre-use-poststerilization-integrity-test)

- Support in risk evaluation
- Expertise in filtration system design equipment characterization
- Dedicated Mlab Collaboration Center to meet with our experts







Choosing the right filters to sterilize your product

Considerations

Filter design

Filtration System Setup

Line Flushing Integrity Testing

Validation

Process Design

Solution

- Maintain a closed system
- Optimum flushing volume (to mitigate E&L risk and adsorption phenomenon)
- Filter integrity test method

- Certification training for operators
- Options to maintain a closed system through <u>flush bags or barrier filter</u>
- Single-use system design expertise to simplify PUPSIT implementation

Pharma & Biopharma Manufacturing & Testing Services





Choosing the right filters to sterilize your product

Considerations

Filter design

Filtration System Setup

Line Flushing Integrity Testing

Validation

Process Design

Solution

Equipment coming in contact with the product must not be

- additive
- reactive
- absorptive

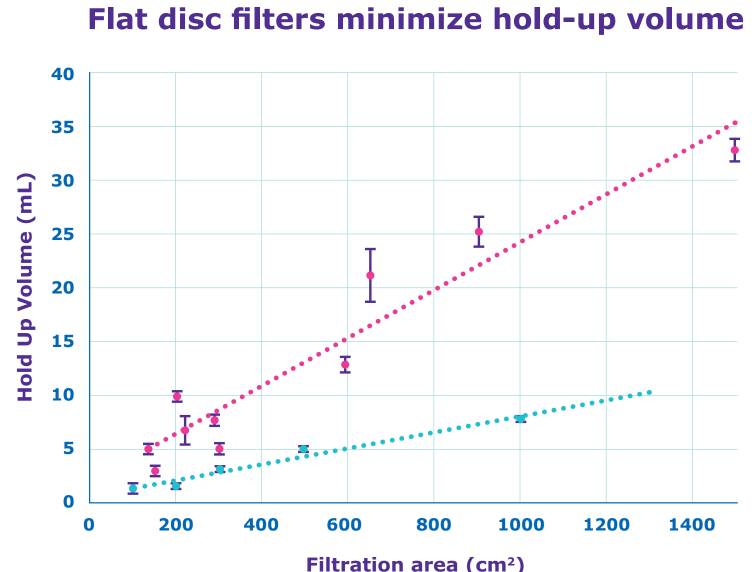
- Validation team experts
 - to help you choose and conduct the appropriate validation services for your aseptic filtration
 - to accelerate and simplify your path to market
- Documentation support to meet your compliance needs

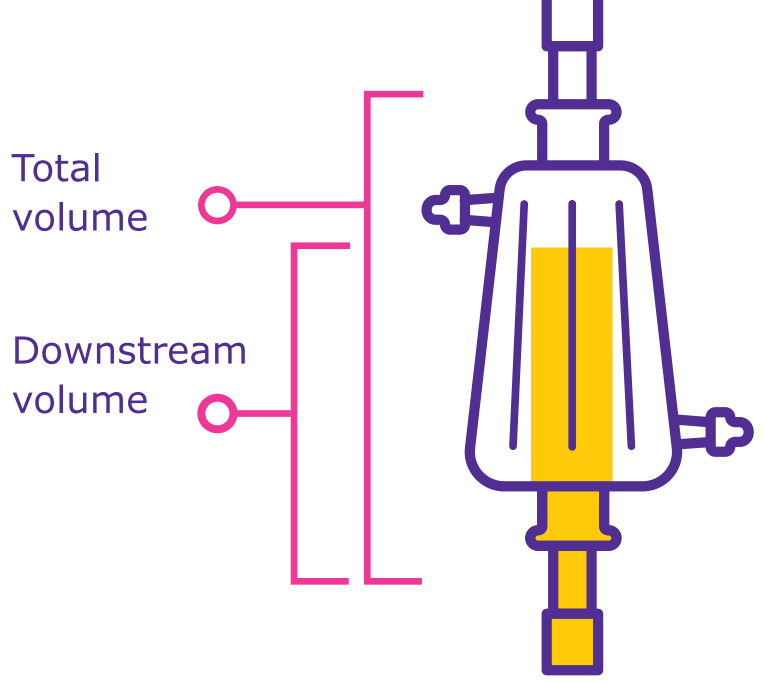
BioReliance® Validation Services

Minimize hold-up volume with stacked disc design



	Filtration area	Total void volume	Downstream volume	Hold up volume
Millipak® FF 20 filter	100 cm ²	38 mL	6.4 mL	1.1 mL
Millipak® FF 200 filter	1,000 cm ²	145 mL	38 mL	7.7 mL
Opticap® XL 2 capsule	900 cm ²	225 mL	30 mL	15 mL

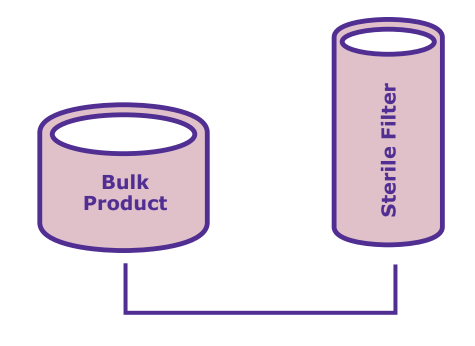








Design options



Filter
Configurations
Single Filter
Sterile Filtration

Single filter ▶

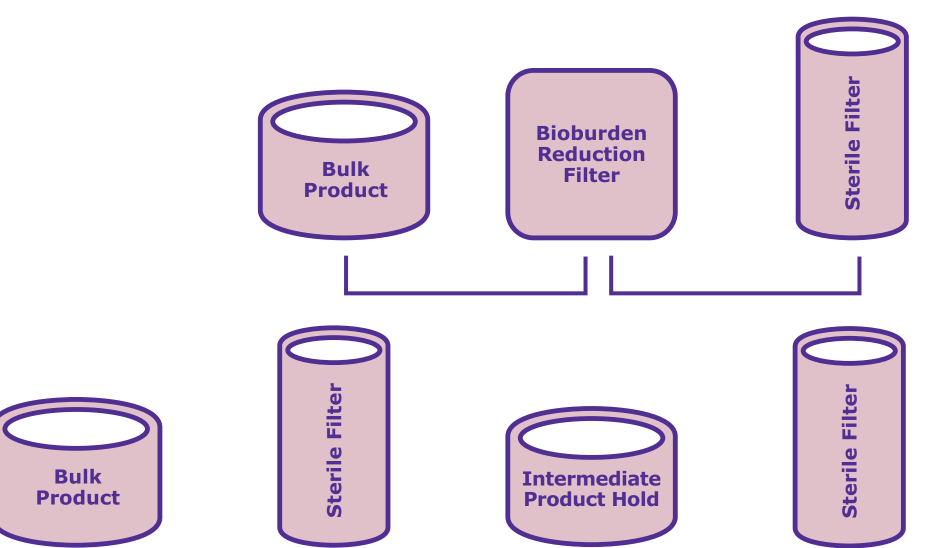
Sterile Filtration

Advantages

- Minimum hold-up volume
- Minimum flushing requirements
- Ease of handling and operation
- Lower filter cost

Disadvantages

- No back-up in the event of primary filter failure
- Filter plugging



Dual FiltersBioburden/Sterile
Filtration

Dual Filters

Filtration

Sterile Filtration

- Hold - Sterile

Dual filter

Bioburden / Sterile Filtration

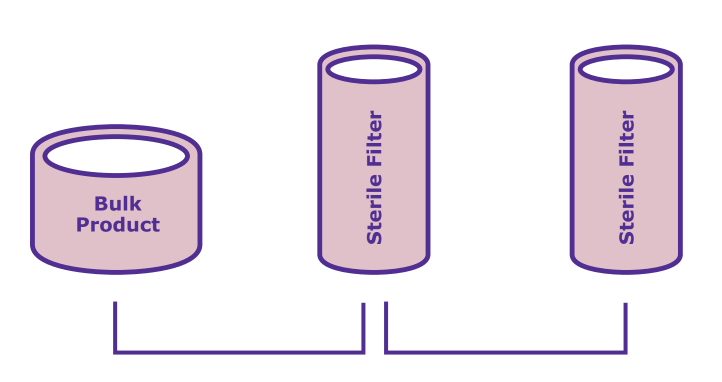
Sterile – Hold - Sterile Filtration

Advantages

- Compliance with regulatory guidance for <10 cfu/100 mL
- Very low plugging risk for primary filter

Disadvantages

- No back-up in the event of primary filter failure
- Higher hold up volume
- Higher cost than single filter
- Higher system complexity than single filter



Redundant Filters Sterile Filtration

Redundant

Sterile Filtration

Advantages

- Compliance with regulatory guidance for <10 cfu/100 mL
- Very low plugging risk for primary filter
- Potential batch recovery if one filter fails IT

Disadvantages

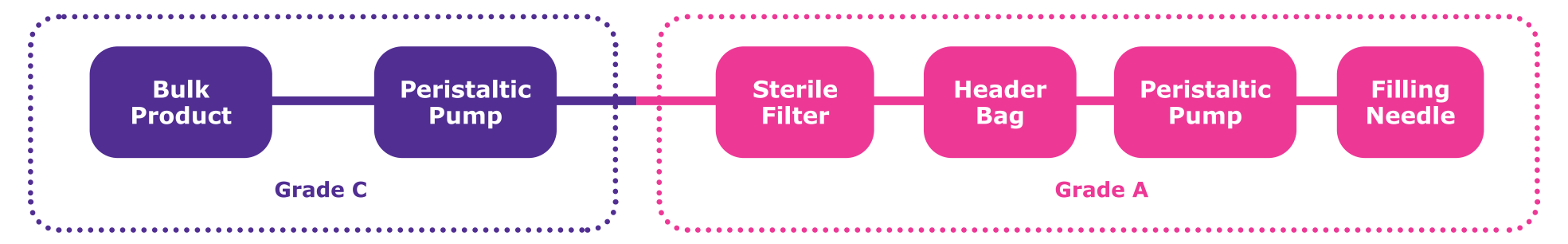
- Higher hold up volume
- Higher cost than single filter
- Higher system complexity than single filter



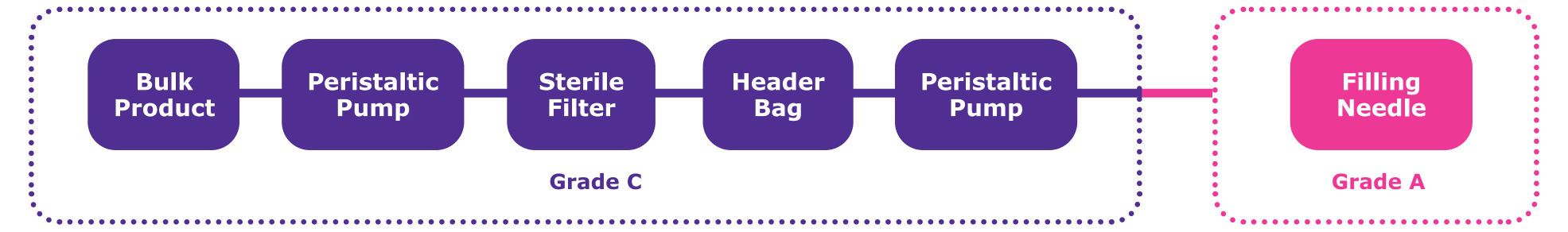
Sterile filter placement

Sterile filter placement considerations: Inside or Outside the Isolator/RABS

Sterile filter in Grade A/ISO 5/Class 100 space



Sterile filter in Grade C/ISO 7/Class 10,000 space



Complete SUS transferred to Grade A

Benefits

 Low risk of fluid path contamination with majority of operations (venting, connections, etc.) occurring inside isolator/RABS

Considerations

• Sterile environment risk if performing open venting

Handling challenges

- Large number of components to transfer from Grade C into Grade A
- Pre-use integrity testing challenge

Only filling needles transferred to Grade A

Benefits

- Majority of operations (instability, flushing, etc.) occur outside isolator/RABS
- Potential to replace filter if integrity test fails

Considerations

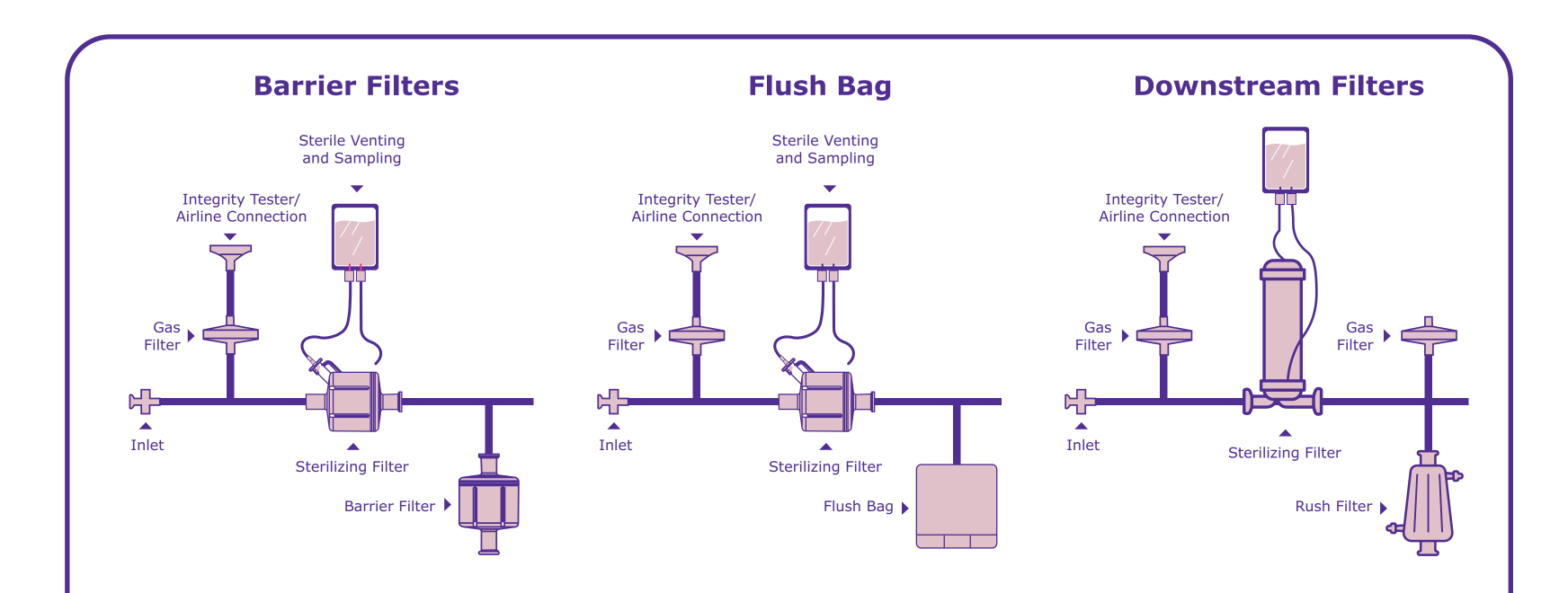
- More potential points for microbial ingress
- Demonstration of assembly integrity more critical

Maintaining a closed system

Details about barrier filter

- Millipak® Barrier filters simplify in-line pre-use, post-sterilization integrity testing (PUPSIT) of single or redundant liquid filtration systems.
- Separate layers of hydrophobic and hydrophilic 0.22 µm Durapore® sterilizing-grade membranes allow the sterile flow of liquid and gas.
- Simplify wetting, flushing and integrity testing of upstream sterile filters in filtration assemblies while removing the constraint of a flush bag or can.





	Barrier Filter	Flush Bag	Downstream Filter
Ability to Retest	Yes	No	Yes
Ability to Dry the Product Filter	Yes	No	Yes
Simple Design	Yes	Yes	No

Filtration

Filling

BioReliance® Validation Services

BioReliance®

Pharma & Biopharma Manufacturing & Testing Services

Validation services for filtration

Extractables

Identify and quantify the extractables which may be extracted out from the filter by employing the Model Solvent Stream Approach and worst case test conditions.

Bacterial retention validation

Validate the performance of your sterilizing grade filter by simulating your product and process conditions. *B. diminuta* is used as the standard challenge microorganism.

Bubble point/diffusion determination

Provide a product-specific bubble point/diffusion value.

Patient safety assessment

Assess the potential impact of the substances that have been detected, identified and quantified on patient safety.



Leachables

Identify and quantify the leachables which leach out from the filter with the use of your actual product under normal processing conditions.

Particle shedding

Provide particle shedding data with the drug product and using actual and scale-down customer worse-case filtration conditions.

Binding study

Show that the filter does not remove unacceptable amounts of stream compounds.

Compatibility study

Assess the chemical compatibility based on key characteristics and provide evidence that the process fluids and conditions do not adversely impact the structure of the filter device.

Click here for more information on the BioReliance® Validation Services



Opticap® Capsules

- Eliminates time and expense associated with stainless steel housings
- Pleated filter membrane



Millipak® Final Fill

- Designed for small volume, high value solutions
- Stacked disc design
- Multi-purpose port that simplifies venting, integrity testing and sampling, and is validated to maintain an aseptic flow path



Millipore Express[®] Membrane

- Polyethersulfone (PES) membranes
- Broad chemical compatibility, pH 1-14
- Exceptionally high flow rates

- Polyvinylidene fluoride (PVDF) membrane
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- High flow rates



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- Verifying the integrity of your filters and processing equipment
- Portable, easy to implement, and automated
- Delivering a simple and intuitive user experience
- Providing optional depth of flexibility to fit your process





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Choosing the right sampling systems to monitor critical parameters across each step

Considerations

Facility Needs Meeting Regulation Validation Ease of Use Requirements **Process Design** Solution Solutions for all process steps Flexibility and speed Safety of operators Wide range of sampling containers, available pre-sterilized and pre-assembled Secure disconnection of sample Preconfigured or configure on site No operator training needed





Choosing the right sampling systems to monitor critical parameters across each step

Considerations

Ease of Use Facility Needs

Meeting Regulation Requirements

Validation

Process Design

Solution

- No need for additional utilities (pressure, steam, ...)
- Option of sampling from Single-Use and Stainless Steel
- Limited investment
- Maintenance & spare parts

- Utilities: only need of crimping tool to disconnect taken sample
- Sampling from Single-Use and Stainless
 Steel tank possible
- No cleaning needs





Choosing the right sampling systems to monitor critical parameters across each step

Considerations

Process Design

Solution

• Right specifications
• Representative samples
• Process risk mitigation
• Accelerate speed to market

• Radility Needs

Meeting Regulation

Foliation

• Process integrity
• No dead leg, flush, or sample dilution
• Closed and validated sampling system
• Sampling actuation evidence and control ▶

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Choosing the right sampling systems to monitor critical parameters across each step

Considerations

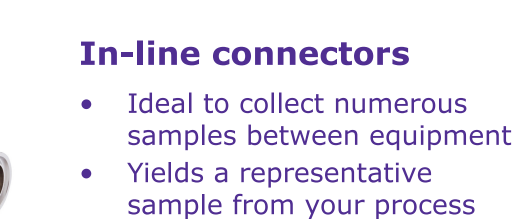
Facility Needs Ease of Use Meeting Regulation Validation Requirements **Process Design** Solution Drug Product shall not be affected by the Validation team experts process conditions and equipment - to help you choose and conduct the appropriate validation services for your sampling system - to accelerate and simplify your path to market Documentation support to meet your compliance needs

Process step options for sampling before sterilizing filtration

BioReliance® Validation Services

Connector for Stainless Steel tank

Emprove® Program



stream

Sampling containers



Bottles

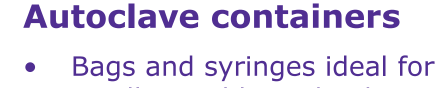
- Bio-neutral plastic bottles ideal for all tests
- Offered in multiple sizes

Without Connector

- Directly at the filtration assembly
- Directly at the Final Fill Filter

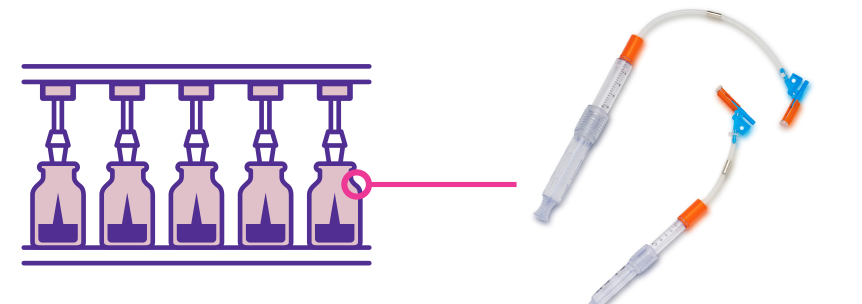






small portable tanks that are fully autoclaved

 Vented to withstand the most aggressive autoclave cycles



Sterile syringes

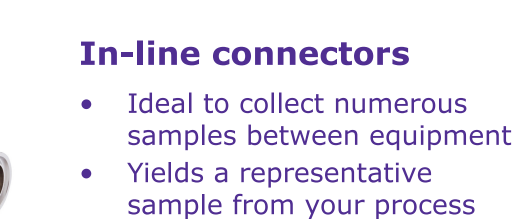
- Contamination free with patented design
- Ideal to save and store high value product

Process step options for sampling before sterilizing filtration

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Connector for Stainless Steel tank

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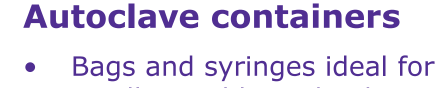
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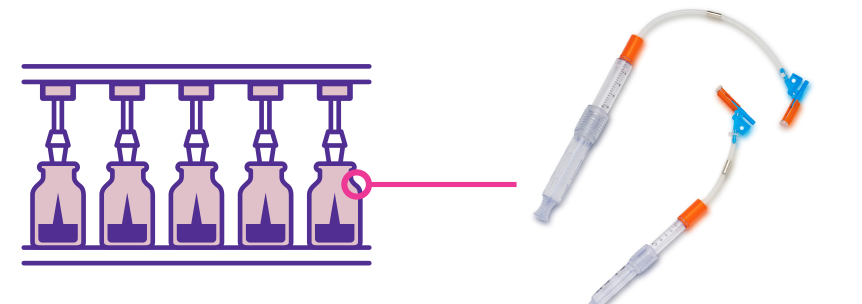






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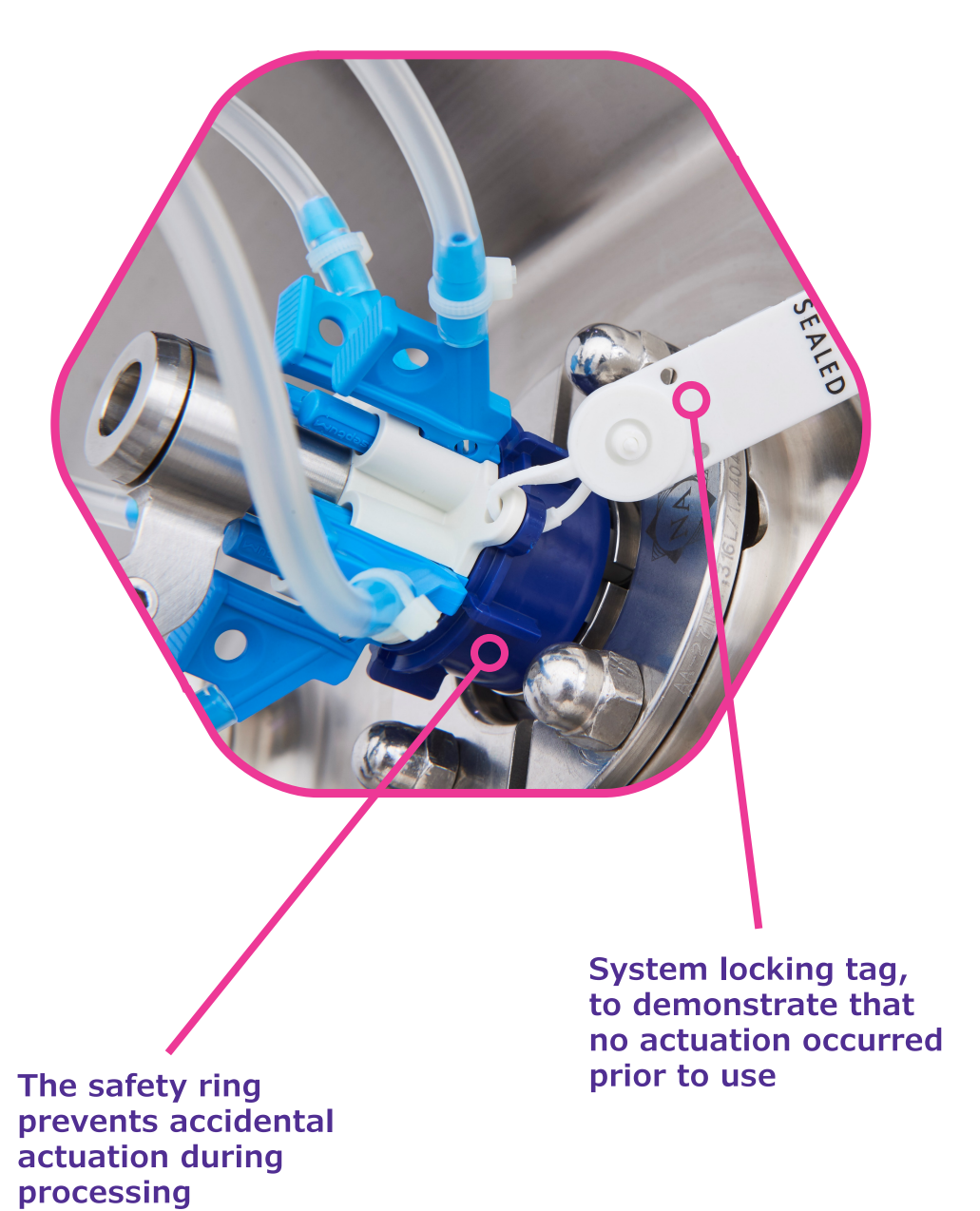
Sterile syringes

- Contamination free with patented design
- Ideal to save and store high value product



Maximum security and reliability

- Eliminating high heat/pressure and glass bottle safety risks
- No steam condensate to dilute sample, avoid requirement for flushing
- "Neutral" containers for accurate testing and storage
- Simple procedure eliminates complexity of training, reducing risk of operator bias
- Eliminating time of steam sterilization and cooling between samples



Validation services for sampling system

Chemical compatibility

Assess the chemical compatibility based on key characteristics, after prolonged exposure with the drug product. Provide evidence that the process fluids and conditions do not adversely impact the structure of the Sampling System

Functionality test

Verify that the Sampling element is within the qualified acceptance criteria after product contact & process conditions simulation.

Non permeation test for bag

Demonstrate the bag is not permeable to disinfectant/ decontamination agent

Click here for more information on the BioReliance® Validation Services

BioReliance® Validation Services

NovaSeptum® GO sterile sampling system

- A fast and easy connecting of the sampling solution to the manufacturing process.
- Available in various designs







Custom



The NovaSeptum® GO high purity bag single and multi-sampling systems are available from 50 to 1000 mL.



The NovaSeptum® GO autoclavable bag single sampling system is available in 50, 100, 250 and 1000 mL.



The NovaSeptum® GO bottle single and multisampling systems are available in 60 to 500 mL.



The NovaSeptum® GO transfer unit is available in different tubing materials (Thermoplastic Silicone, and C-Flex®).



The NovaSeptum® GO AV single and multisampling units are available from 5 mL and 20 mL syringe sizes.

BioReliance® Validation Services

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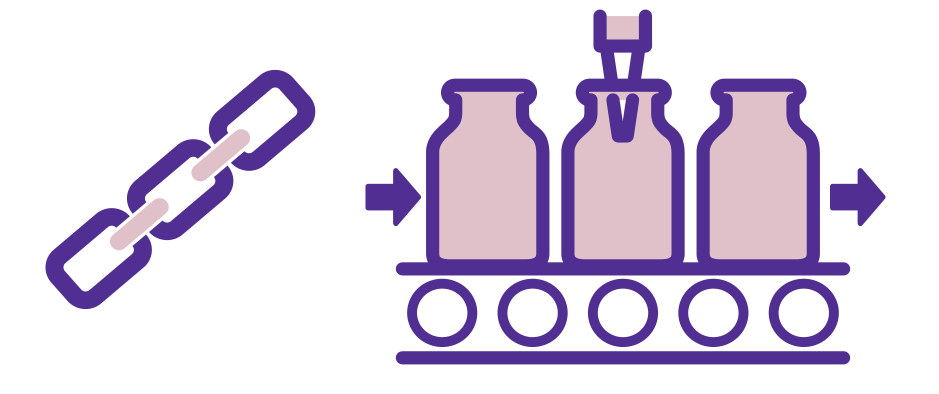
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Key steps for filling

Sterile connection •

- Robust design delivers consistent, reliable performance
- Completely closed flow path 100% air-integrity tested in manufacturing



Final fill assemblies >

- Extensive library of prequalified components providing flexibility for customized assembly design
- Easy integration with different filling machines
- Extensive technical support in design, risk assessment and testing services





Chosing the right sterile connectors to maintain an aseptic fluid path

Considerations

Sterility Assurance Handling **Application Validation Process Design** Solution Robust design delivers consistent, reliable Quick and easy connectivity to avoid operator mistakes performance





Chosing the right sterile connectors to maintain an aseptic fluid path

Considerations

Validation Handling **Sterility Assurance Application Process Design** Solution Completely closed flow path 100% air-integrity tested in manufacturing > • Sterile connection could be made in both Aerosolized microbial challenge test – non-classified and classified area worst case simulation for validation Eliminate contamination risk associated with aseptic connections





Chosing the right sterile connectors to maintain an aseptic fluid path

Considerations

Application Sterility Assurance Validation Handling **Process Design** Solution Sterile fluid transfer Sterile to sterile connection Connect sterilized fluid paths, integrate Steamable hard-piped connection to steamable hard-piped process equipment disposable flow path with disposable sterile fluid paths Multiple connections/disconnections Single and multiple actuations

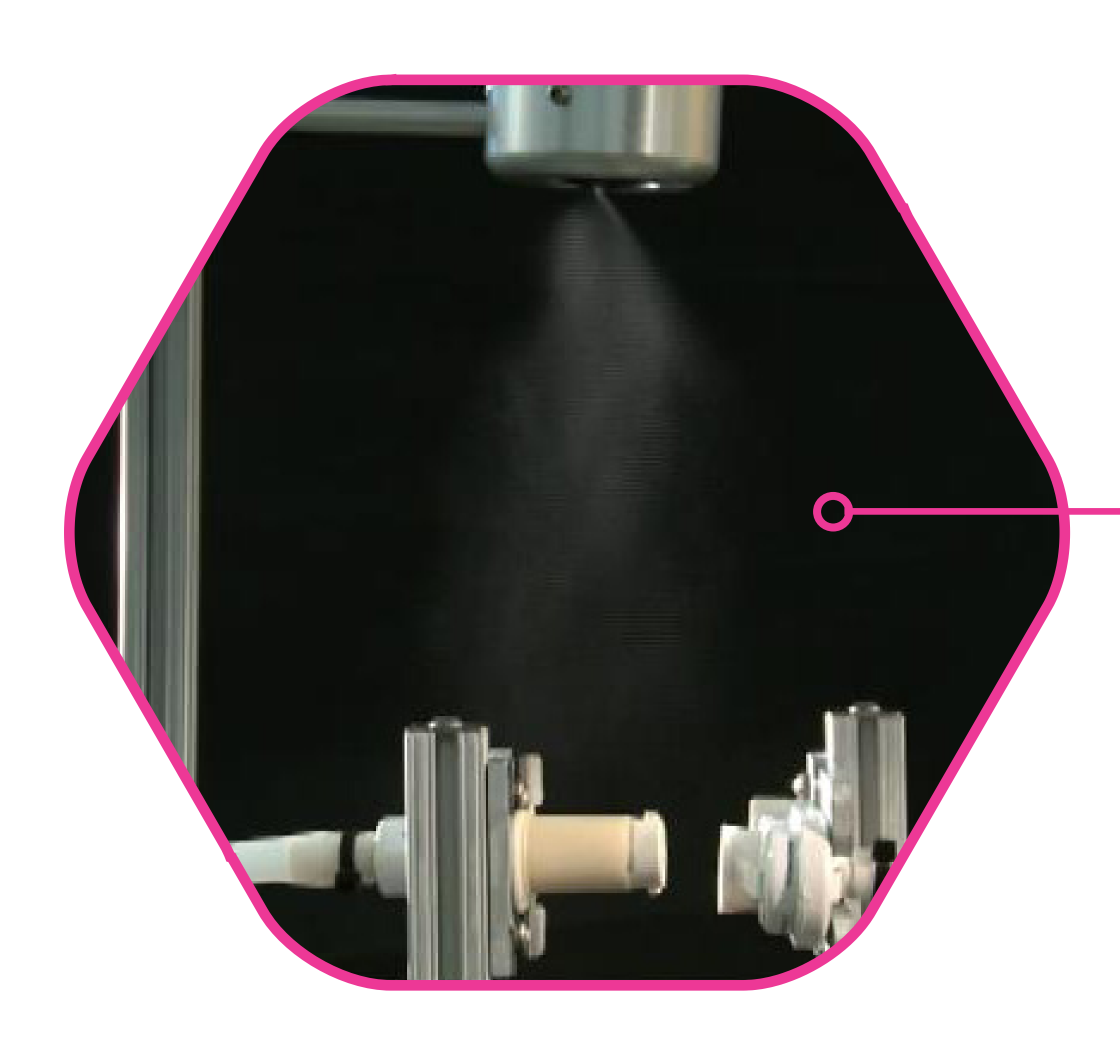
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Chosing the right sterile connectors to maintain an aseptic fluid path

Considerations

Sterility Assurance Handling **Application Validation Process Design** Solution Drug Product shall not be affected by the Validation team experts process conditions and equipment - to help you choose and conduct the appropriate validation services for sterile connectors - to accelerate and simplify your path to market Documentation support to meet your compliance needs



Brevundimonas diminuta, >4 x 10⁶ cfu per connector set

Sterility assurance during actuation

Two tests were conducted to challenge the ability of Lynx® S2S connectors to prevent ingress of microorganisms in a non-classified area.

Tests were performed to confirm that the Lynx® S2S connectors showed no ingress of the challenge organism, *B. diminuta*.

Bacterial aerosol

The connector was connected and actuated in the presence of a bacterial aerosol at a minimum concentration of 1×10^6 cfu/connector.

Direct bacterial soiling

B. diminuta was applied to the mating surfaces of the male and female Lynx® S2S connector lumen plugs, followed after a setting time, by connection and actuation.

All Lynx® S2S connectors met the acceptance criteria for the Bacterial Aerosol Test and the Direct Bacterial Soiling test, assuring a sterile connection can be made in non-classified areas.

Validation services for sterile connectors

Chemical compatibility

Assess the chemical compatibility based on key characteristics, after prolonged exposure with the drug product. Provide evidence that the process fluids and conditions do not adversely impact the structure of the connectors.

Extractables

Identify and quantify
the extractables which may
be extracted out from the
connectors by employing the
Model Solvent Stream Approach
and worst case test conditions.
Analytical methods used are
NVR, TOC, FTIR, RP-HPLC and
GC-MS (when applicable).

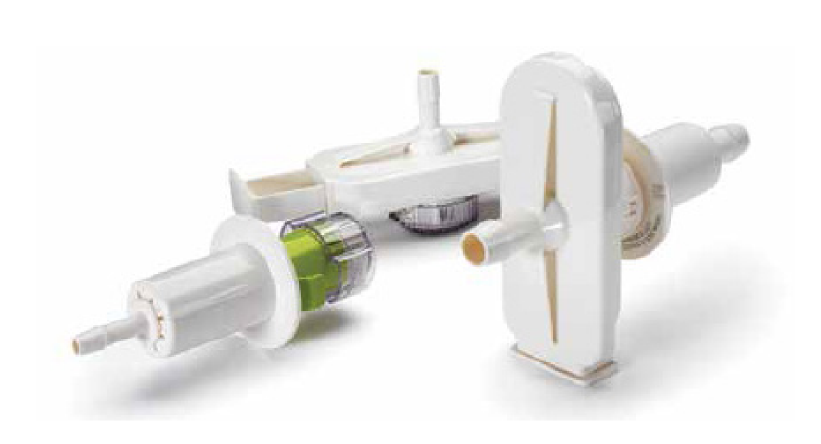
Patient safety

Assess the potential impact of the substances that have been detected, identified and quantified on patient safety. This assessment is done by a toxicologist.

Click here for more information on the BioReliance® Validation Services •

BioReliance® Validation Services

Lynx® product family







Lynx® S2S

Sterile to Sterile Connector

A single actuation, disposable device for connecting sterilized disposable flow paths

Lynx[®] ST

Steam-To Connector

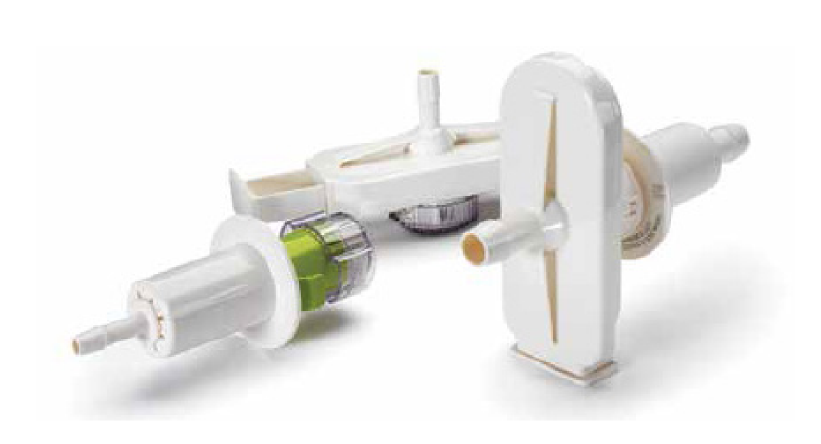
Designed to connect steamable hard-piped processing systems to sterilized disposable flow paths

Lynx® CDR

Connect, Disconnect, Reconnect

BioReliance® Validation Services

Lynx® product family







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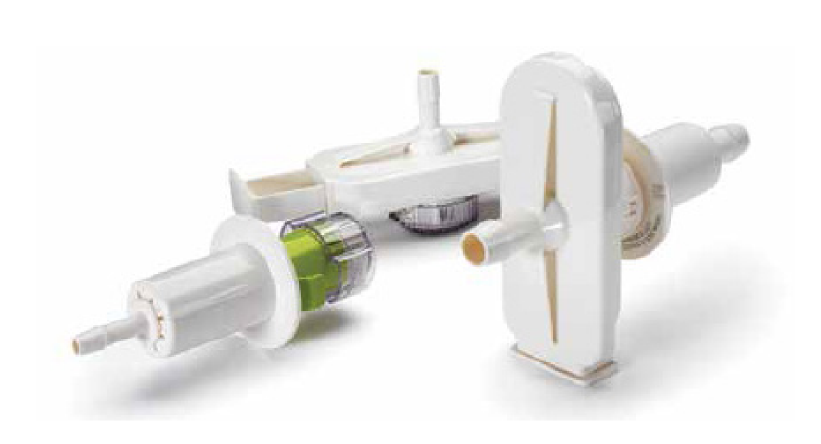
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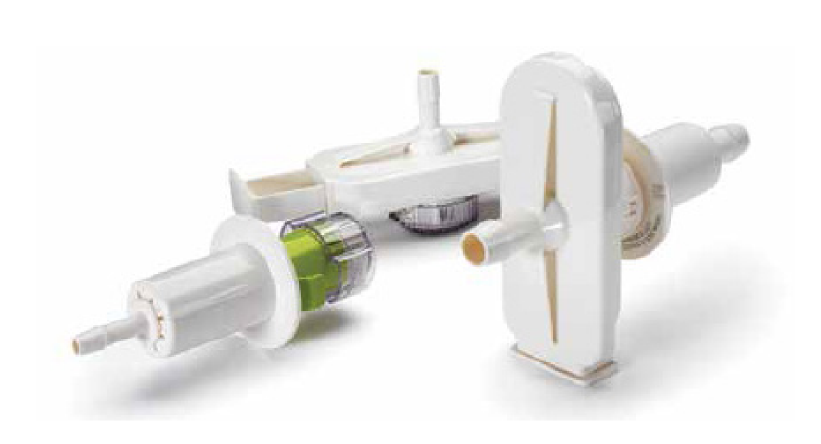
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Filling assemblies are designed to fit for purpose

Considerations **Risk Mitigation Validation Customization** Quality **Enhanced Economics Process Design** Solution • Extensive single-use component-library Full single-use system Integrating into various filling machines Pre-qualified components Integrating with isolator / RABS Flexibility for customized assembly design (restricted access barrier system) filling Expertise to collaborate with machine machines vendors, to design, integrate and validate single-use assemblies for your filling process





Filling assemblies are designed to fit for purpose

Considerations **Customization Validation Risk Mitigation** Quality **Enhanced Economics Process Design** Solution Expertise to refine system design Components placement (pump tubing, header bag, filter) inside/outside of Designed and packaged for ease-of-use isolator On-site support during machine Operator's handling and installation qualification steps challenges





Filling assemblies are designed to fit for purpose Considerations **Risk Mitigation Customization Validation Enhanced** Quality **Economics Process Design** Solution No particle risks • Fully sterile assemblies, manufactured in dedicated clean room areas Sterility assurance Various material and release testing • Integrity Testing and claims >

Pharma & Biopharma Manufacturing & Testing Services



Filling assemblies are designed to fit for purpose

Considerations **Customization Risk Mitigation Validation** Quality **Enhanced Economics Process Design** Solution Drug Product shall not be affected by the Validation team experts process conditions and equipment - to help you choose and conduct the appropriate validation services for your final fill assemblies - to accelerate and simplify your path to market Documentation support to meet your compliance needs





Filling assemblies are designed to fit for purpose

Considerations **Risk Mitigation Customization Validation** Quality **Enhanced Economics Process Design** Solution • Increase efficiency and minimize Partnership from initial specification to optimization to validation of your singledowntime use system Cost effectiveness Reduce cleaning time



Material and release testing claims

Criteria:

Gamma compatibility >40kGy

Functional testing

Regulatory statements (Animal Origin, Latex, BPA, etc.)

USP <88> Class VI

USP <85> Endotoxin

USP <788> Particulates

USP <661> Physiochemical

Shelf life > 2 years

Sterility per ANSI/AAMI/ISO 11737

Bacteriostasis/Fungistasis

Bioburden

Mobius® assembly leak testing

- Gold certified assemblies are 100% tested
- Testing performed using pressure decay method

Packaging and transport validation

Mobius® packaging protects assemblies from the rigors of typical shipping conditions. Representative Mobius® Final Fill assemblies have been proven to be integral after being subjected to a packaging validation per the International Safe Transit Association (ISTA) 2A Procedure.

Validation service for final fill assemblies

Chemical compatibility

Assess the chemical compatibility based on key characteristics, after prolonged exposure with the drug product. Provide evidence that the process fluids and conditions do not adversely impact the structure of the connectors.

Extractables

Identify and quantify
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be extracted out from the
connectors by employing the
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GC-MS (when applicable).

Patient safety

Assess the potential impact of the substances that have been detected, identified and quantified on patient safety. This assessment is done by a toxicologist.

Click here for more information on the BioReliance® Validation Services •



Efficient and cost effective

Customer case study:

A filling campaign setup, installation and production time was reduced from 36 to 12 hours by using a Mobius® single-use final fill solution versus a traditional stainless steel system.

Features and Benefits:

- Reduced upfront capital investment
- Reduced risk of cross contamination and enhanced operator safety
- Flexibility and increased filling productivity
- Advanced single-use filtration technologies to maximize yields

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BioReliance® Validation Services

Assembly product page

Example of a final fill assembly:

In close collaboration, discussion and review with our technical experts, final fill assemblies are full custom designed, depending on your filling machine design and process needs.

- Header bag design and size
- Tubing lengths and materials
- Flush bag / barrier filter
- Various types of sterile connectors
- Various filling needle styles
- RTP (rapid transfer port) bag and tri-clamp manifold pass-through
- Unique needle packaging and tubing management solution



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Your fast track through regulatory challenges

Example of a final fill assembly:

High-quality products, comprehensive documentation, and superior customer support facilitate your efforts in qualification, risk assessment, and process optimization.

Confidently speed your way through the regulatory maze and fast track your new drug to market.

Emprove® Dossiers for Chemicals

Emprove® Dossiers for Chemicals provide three levels of Information Supporting Quality Risk Assessment and Regulatory Requirements

Emprove® Chemicals Portfolio:Starting and Raw Materials

Simplifying the selection process

Our Emprove® Chemicals portfolio contains over 400 raw materials. To address different levels of risk, and to simplify and streamline the selection process, the Portfolio is divided into four categories: Emprove® Evolve, Emprove® Essential, Emprove® Expert, Emprove® API

Emprove® Dossiers for Filtration and Single-Use

Emprove® Dossiers for Filters are grouped according to product families with the same materials of construction, production processes, and packaging components.

Emprove® Dossiers for Single-Use Components are available beginning with the most commonly used components in the process. Assembly level information can be developed from the individual dossiers available for components forming the part of the single use assemblies.



Emprove® Dossiers for chemicals



Material Qualification Dossier

Supports raw material qualification and speeds up drug filing preparation.

In line with CTD chapter 3 quality (adapted for excipients)

- General information
- Manufacturer
- Characterization
- Control of drug substance
- Reference standard
- Materials
- Container closure system
- Stability summary

Emprove® Suite Subscription

Unlock Full Access to all dossiers today with the Emprove® Suite Subscription

- 1, 2, or 5-year subscriptions
- 24/7 full online access
- Constantly updated
- Optimized searches



Management Dossier Quality Management Dossier

Answers questions during quality risk assessment according to ICH Q9 and EU 2015/C95/02.

- Supply chain information
- Product quality self-assessment
- Audit report summary
- Stability data**

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**Available for Emprove® Evolve™, Emprove® Essential and Emprove® Expert products only



Operational Excellence Dossier

Supports process optimization and safety risk assessment activities.

- Elemental impurities information
- Product quality report
- Analytical procedure
- Technically Unavoidable Particle Profile (if applicable)



Emprove® Dossiers for filtration and single-use

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Emprove® Quality Management Dossier



Material Qualification Dossier

Supports product qualification and speeds up regulatory filing preparation. Includes content on the manufacturing process, product specifications and various qualification criteria (product validation data), regulatory statements, and more.

- General information
- Manufacturing flow chart
- Product validation and qualification
- Specifications (design and release criteria)
- Materials of construction
- Extractables summary**
- Regulatory statements (Animal origin, e.g. Animal Origin, BPA etc.)

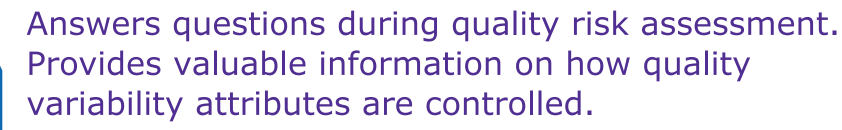
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Quality Management Dossier



- Quality Self-assessments
- Chain of custody
- Supplier and CMO management
- Shelf life testing and results
- Packaging and Sterilization Validation

Emprove® Operational Excellence Dossier

Operational Excellence Dossier

Support process optimization and safety risk assessment with detailed extractables profile (BPOG extractables protocol and USP <665> draft chapters and information on elemental impurities (ICH Q3D)).

- Extractables Report**
- Elemental impurities summary
- Analytical procedure



Emprove® chemicals portfolio

Starting and Raw Materials

Emprove® Evolve

For early stage or (bio-)pharmaceutical manufacturing

Fills the gap between lab-grade and GMP compliant raw and starting materials. This product line provides detailed and transparent supply chain information and documentation to support risk assessments for critical raw materials used in manufacturing processes.

Emprove® Essential

Moderate Risk Applications

Designed for moderate risk applications, Emprove® Essential products offer compliance to IPEC PQG GMP Guide and/or EXiPACT™ Certification Standard, supply chain transparency and regulatory support designed to assist drug manufacturers' formalized risk assessments. They are produced according to controlled manufacturing processes. Critical parameters such as elemental impurities and residual solvents are characterized by using validated analytical techniques.

Emprove® Expert

High Risk Applications

Addresses higher risk applications where the lowest microbiological and endotoxin levels are of utmost importance. Along with the risk management features of Emprove® Essential, the Emprove® Expert line goes even further: The cGMP manufacturing processes are designed to yield products with specified low microbiological and endotoxin levels, thus supporting the overall risk mitigation strategy.

Emprove® API

Support Final Drug Product Compliance with International Standards

Manufactured in Europe to meet the quality and regulatory requirements of active pharmaceutical ingredients, according to ICH Q7 GMP. In order to support final drug product compliance with international standards, our Regulatory Affairs team offers dedicated support with access to extensive documentation including DMFs, CEP and ASMF.

Details to each test >

Extractables evaluation*

Leachables testing

Patient safety evaluation*

Bacterial retention testing*

Integrity testing*

(Diffusion, Bubble Point, Rinsing)

Compatibility evaluation*

Binding studies



^{*}Validation Requirements for Regulatory Submissions

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Extractables

Identify and quantify the extractables which may be extracted out from the filter by employing the Model Solvent Stream Approach and worst case test conditions.

Leachables

Identify and quantify the leachables which may be leached out from the filter with the use of your actual product under normal processing conditions.

Patient safety assessment

Assess the potential impact of the substances that have been detected, identified and quantified on patient safety.

Bacterial retention validation

Validate the performance of your sterilizing grade filter by simulating your product and process conditions. *B. diminuta* is used as the standard challenge microorganism.

Bubble point / diffusion determination

Provide a product-specific bubble point/diffusion value.

Compatibility study

Assess the chemical compatibility based on key characteristics and provide evidence that the process fluids and conditions do not adversely impact the structure of the filter device.

Binding study

Show the filter does not remove unacceptable amounts of stream compounds.

Particle shedding

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EMDMillipore.com

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