All about Pre-filled Syringe Systems From Initial Development to Final Fill Finish

Plunger, Needle Shield, Tip Cap Christa Jansen-Otten

Gothenburg, October 20th 2023







Agenda – DAY 2

The "Ready-to Fill" Syringe

Material • Shape • Properties • Siliconization • Impact of different drug • Nest and Tub • Needles and LL • backstops • Rods • Regulatory Guidelines

Plunger Stoppers, Needle Shields, Tip Caps Materials • Properties • Functionality • Production • Regulatory

Manufacturing Aspects in Fill & Finish and Assembly Bulk versus Nested • Nest Sizes • Rod insertion • Handling of Syringes, Labeling • Glass to Glass Contact

Assembly of Syringes and Administration Devices Pen Injectors • Safety systems • Autoinjectors • Manual vs Automated

Design Independent Assembly

Hands-on Session 2, Mind map, Lottery





Customer Impact - Demands on Packaging Components are Increasing



- Particulate reduction/foreign matter
- S Concerns regarding extractables/leachables
 - Ultra-clean components needed
 - New ways to deliver medicine
- Functional performance of components
 - High-speed lines
 - Complex devices
- Moisture Vapor Transmission Rate





- S Container closure integrity (CCI)
- New manufacturing approach
 - Flexibility
 - Time to market
 - Total cost of ownership (TCO) focused
- S Functional performance of components
 - High-speed lines
 - Complex devices





Considerations in Selection of PFS Components

COMPATIBILITY WITH DRUG	APPLICATION – MANUAL OR AUTO SYSTEM	CONTAINER CLOSURE INTEGRIETY (CCI)	QUALITY SPECIFICATIONS	FINISHIING
 Type of drug pH Viscosity Excipients 	 Break loose & glide force requirements Accuracy of delivery volume 	 Interference fit of plunger with barrel Sealing ribs and their function Preservation of drug potency and sterility 	 Particulate level – visible & sub-visible Dimensional control Endotoxin level Bioburden level Visual defects 	 Mode of sterilization Lubricity Consistency







Rubber material







Why Use a Rubber Material?



CONNECTING PEOPLE SCIENCE AND REGULATION®

pda.org

Polymers: thermosets and thermoplastics

PDA*







Main Elastomer Types Used for Parenteral Applications

Natural Rubber (NR) – from Hevea Brasiliensis	Nitrile Rubber (NBR)
Isoprene Rubber (IR) – synthetic equivalent to NR	Ethylene-Propylene Rubber (EPM/EPDM)
Styrene-Butadiene-Rubber (SBR)	Isobutylene Isoprene Rubber (IIR, Butyl Rubber)
Butadiene Rubber (BR)	Halogenated Butyl Rubber (XIIR) – Br, Cl





Elastomers Closures General Composition



Additives

can be curing agents, antioxidants, accelerators, activators, protective agents, colorants, plasticizers, acid scavengers, light and heat stabilizers, lubricants, anti-static agents, etc. Approximate Composition of an Elastomer Component



Fillers

are mainly defining the physical properties

 \rightarrow "Ask your supplier for potential extractable lists"





Elastomeric Formulations for Pharmaceutical Use - Properties Butyls/Halobutyls



¹extractables & leachables ²design dependent

10 COPYRIGHT © PDA



pda.org

pda.org

PDA*

Elastomeric Formulations for Pharmaceutical Use - Properties synthetic Polyisoprene







Potential Issues: Needle Shields and Tip Caps









Supporting Documents





pda.org



Supporting Documents: Example

- Technical drawings
- Formulation Characteristics
- Elastomer Formulation Biocompatibility
- Technical Bulletins and Reports
- Theoretical Material Extractable List
- VeriSure[®] Extractable Technical Package
- Material Characterization Package
- Regulatory Compliance Bulletins
- Product Specifications
- DMF
- Certificates











Formulation Documents: Example

VVG5					
	T			Version: Supersedes:	Revision 6 Revision 5
	FOR	ULATION CHAR	CTERISTICS		
W	VEST FOR	RMULATIO	N 4023/50	GRAY	
Note: The formulations lis	sted above conform to	o the specifications and pro	perfies represented be	elow as an exampl	le.
1. FORMULATION SP	ECIFICATIONS				
The following tests and sp specific gravity specification	ecifications may be u	sed to confirm the compos	tion of the base elastor JuroTec® or LvoTec™	meric formulation.	The ash and
beome Branny sheemoone	no noted do not debi	is crossies man remotes,	1010000010100	oounigo.	
	Test	Inite	Specification]	Method
	Test	[Units]	Specification	[Method
	Test	[Units]	Specification		Method
2. GENERAL DESCRIF	Test	[Units]	Specification	[Method
2. GENERAL DESCRIF The following is a ger	Test PTION heral description of the	Units	Specification		Method
2. GENERAL DESCRIF The following is a ger Elastorrer Type:	Test PTION heral description of th	Units	Specification		Method





pda.org

Formulation Documents: Example

		Rev. 5
N	lest Item: 4023/50 Grey	
	TABLE OF CONTENTS	
1	INTRODUCTION	2
2	ANIMAL DERIVED MATERIALS	2
3	HEAVY METALS	2
4	NATURAL RUBBER LATEX/DRY NATURAL RUBBER CONTE	ENT 3
5	ADDITIVES OF CONCERN	
6	ROHS	4
7	FOOD REGULATIONS	
8	PROPOSITION 65	
9	ELEMENTAL IMPURITIES	
10	HALAL	6
11	KOSHER	6



Theoretical Material Extractables List: 4023/50 Gray

Below is a summary of the potential chemical entities that could be extracted from West elastomer formulation 4023/50 Gray based on the materials that are used in the formulation. Since each drug application is unique, it is possible to form new reaction products from the closure or from a combination of the closure and the drug product components.



pda.org



Uncontrolled Drawings: Example









Rigid Needle Shields and Tip Cap





pda.org

pda.org



Pre-filled Head Designs ISO 11040-4 require different closure design solutions



Head design of glass barrel with a 6% Luer cone



Head design of glass barrel with a 6% Luer cone for Luer Lock (LL)



Head design of glass barrel with staked needle











West Rigid Needle Shields



Rigid Needle Shields [RNS] are a safe & efficient closuring system for Prefilled Syringes with staked needles





Design examples of Rigid Needle Shields

RNS 1/2" [13 mm]

Needle length used for subcutaneous drug injection (into the tissue layer between the skin and the muscle)



RNS 5/8" [16 mm]

Needle length used for intramuscular drug injection (deep into the muscles)



pda.org



Solution

0	

Designed for existing assembly machine and filling equipment.



Fits to ISO Norm 11040-4 glass syringe with staked needle

Suitable also for polymer (e.g. COP) syringe





High gas permeation rubber formulation combined with sterilization windows of the rigid shell allowing effective sterilization by ethylene oxide or steam





Advantages of Rigid Needle Shields vs Soft Needle Shields



Rigid Needle Shields are the preferred closure for staked needle syringes





Example of various Tip Caps for Luer and Luer Lock Syringe



Tip Cap to be inserted a rigid plastic cap # 3155



Easy Turn Tip Cap # 3131



Mushroom Rip Cap # 3379



Multiple rubber formulation options (halobutyl and synthetic isoprenes)





pda.org

Barrier Film & Coatings







Films and Coating Technologies



- Film sheet (e.g., PTFE, ETFE) that is laminated to elastomeric component during the molding process
 - - Barrier function, e.g., FluroTec[™] film
- Coating liquid or vapor that is sprayed, tumbled or vapor deposited onto the elastomeric component
 - Lubricity, e.g., B2-Coating
 - Lubricity and barrier function





Film properties



The blue color in**dicates** FluroTec™ film



Structure of Poly(ethylene tetrafluoroethylene) (ETFE)

	FluroTec [™] film is
\bigcirc	based on poly(ethylene tetrafluoroethylene)
•	smooth surface
•	very adherent to elastomers (either bromo- or chloro-butyl)
•	translucent
•	compatible with sterilization by either: • autoclave • gamma irradiation
•	Applied during the compression molding process





Film has a low level of Interaction

ETFE acting as a barrier reduces transport in two directions



	FluroTec™ film
$\overline{\mathbf{O}}$	No reactive functional groups
•	chemically inert – mitigates chemical migration
•	resistant to degradation
•	supports reduction in absorbance

Very Low Surface Energy → Very Low Level of Interaction!





Fluoropolymer film coating Significantly Reduce Leachables



The drawn blue line indicates an estimated identification threshold of 0.5 µg/unit, which is below the Product Quality Research Institute recommended safety concern threshold for parenteral drug products

Non-laminated elastomers showed approximately eight volatile organic compound (VOC) peaks estimated to be > 0.5 µg/unit

Elastomers with FluroTec[™] film did not show any peaks > 0.5 µg/unit [blue line]

Most marketed biopharmaceuticals use fluorpolymer-coated component technology (FluroTec[®] film)

29 COPYRIGHT © PDA

West Technical Report 2019/210 FluroTec® Protecting Drug Product Quality Safety



pda.org







Lubricity Coating: Classical Silicone Oil

<u>Polydimethylsiloxane</u> *DuPont* [™]*Liveo* [™] 360 *Medical Fluid** added during washing operation into the washing drum:

- 350 centistokes \rightarrow USA
- 1000 centistokes → Europe



* <u>Example</u> of silicone oil used by West







Plungers





pda.org



Facilitating Life Cycle: Seamless Transition from Vial to Prefilled Syringe format





Main requirements for Prefillable Syringes Plungers

Delivers a smooth injection profile [break loose & glide forces profile]

Compatibility with the drug product

Compatible with gamma-irradiation and final steam sterilization treatment

Compatible with glass and plastic (COC/COP) barrels

Good compression set properties

Maintains Container closure Integrity

Well performance on fill-finish equipment

Optimized Break Loose & Extrusion Profile

Low Part-to-Part Variability







Plunger ISO 11040-5



b) Plunger stopper with thread (PST)



a) Plunger stopper with snap lid (PSL)

Nominal inner diameter	Nominal volume	Туре	d ₁ a		d2 ^a		d3 ^a		h ₁ a		h2 ^a	
d2b	ml		nom.	tol.	nom.	tol.	nom.	tol.	nom.	tol.	nom.	tol.
4,65 ± 0,1	0,5	PSL	5,2 to 5,3		4,1 to 4,2		2,5		6,85 to 7,0	- ±0,4	5,3	±0,35
$\textbf{6,35} \pm \textbf{0,1}$	1 (long)		6,8 to 7		5,9 to 6		2,6	±0,2	7,65 to 7,85		4,5	±0,3
8,65 ± 0,2	1 to 3		9,05 to 9,25	±0,1	7,6 to 8		4,7		7,7 to 7,85		4	
$11,85\pm0,2$	5	PST	12,5 to 12,7		10,5 to 11,15	±0,15	5,2 to 5,6		8,5		6,0	
$\textbf{14,25} \pm \textbf{0,2}$	10		15 to 15,3		13,5 to 13,75		7,4 to 7,6		8,5 to 10		6 to 6,2	
9,05 ± 0,2	20		19,9 to ±0,1 20,1	±0,15	18,4 to 18,6		10,7		13,45 to 13,50		7	

West standard components are compatible with ISO glass barrels



pda.org





Example of Prefillable Syringe Plungers - Portfolio at West






37 COPYRIGHT © PDA

Examples of Prefilled Syringe plunger designs







Manual PFS to Auto-injector Challenges







Syringe functionality with high variability





Auto-injector Reliability Risks

If injection times vary between doses with an autoinjector:

- > Patient may stop dose if too long
- > Patient may question quality of the product



Critical design factors must be considered, especially functional compatibility

- > Break lose and glide forces (max/min)
- Spring falling rate forces (max/min)









Performance: two different laminated 1 ml long Plungers



40 COPYRIGHT © PDA

Source: West TR 2013/147





Break-loose and Glide Force - 1 ml Long Plungers - Curves represent averages of 30 plots – Example -



- Break-loose forces are on average 50% lower for NovaPure[®] plungers at all viscosities and all timepoints
- Less variability over time with NovaPure[®] plungers especially for high viscosities

Study Extract: TR 2018/191





Evaluation of Plunger Movement During Transit Conditions- Example

West Plungers Evaluated: 1-3 ml FluroTec[™] film laminated plunger and 1-3 mL NovaPure[®] plunger

- Headspace Values
 - 6 mm (exaggerated vent-tube placement)
 - 3 mm (typical vent-tube placement)
 - <1 mm (typical vacuum placement)



Altitude	Significance
8,000 ft	Pressurized Jet
12,000 ft	Mountain Passes
16,000 ft	Unpressurized Jet
20,000 ft	Highest Cargo Jet Altitude on Record

- Linear correlation between pressure and movement
- Higher headspace volume leads to stronger movement
- NovaPure[®] and legacy plunger performance is comparable





Processing











Elastomer Manufacturing Process

Raw Material Weigh Up

- Formulation control SAP
- Only approved raw materials
- Electronic weigh check against the ingredients list

Compounding (Mixer/Open Mill)

- Distribute ingredients uniformly throughout the polymer matrix
- Use shear to reduce the molecular weight of the polymer and allow the ingredients to disperse



Internal Mixer



Open Mill





Mixing Control (Mill Control)

Curing of ISO – standard sample for testing purposes				
specific gravity	per batch			
Shore A of vulcanized sample	per batch			
dispersion of vulcanized sample	per batch			
color of vulcanized sample	per batch			
ash content	every 10 th batch plus 1 st and last			
rheology of the compound	every 5 th batch plus 1 st and last			



Vulcanized Test buttons







Elastomer Manufacturing Process

Batch Off or Calendaring

• Intermediate step that allows the compounding facility to hold or distribute rubber stock prior to extrusion



Extrusion (Calendar/Dispersion Enhancement System)

- Aids in reducing undispersed materials
- Form the compounded rubber into panel shape, required for compression molding





pda.org

Pharmaceutical Rubber Manufacturing

Different 'shapes' need different molding technology:





pda.org



Elastomer Manufacturing Process

Compression Molding

- Mechanical force creates the shape
- Heat forms crosslinks and imparts final physical properties to the part







Elastomer Manufacturing Process

B2-Coating Application

• Applied to the top and/or bottom of the molded panels

Trimming

• Parts are trimmed from the molded panels







Elastomer Manufacturing Process

Rinse

- Removes Processing Aids
- Not a pharmaceutical wash



Pharmaceutical Wash Process

- Pharmaceutical wash process for Ready-to-Sterilize (RS) product
- Application of silicone (if applicable)







Pharmaceutical Wash Process

- 0
- Validated process according to GMP to demonstrate an endotoxin content reduction by at least 99.9% (3.0 log₁₀).
- Components are unloaded from the washer in a Zone 5 clean room
 - All associated process data is filed in Drug Master Files (DMF) with FDA and Health Canada.
 - Particulate, bioburden and endotoxin are reported in the quality certificate provided with every batch



NOVAPURE Westar SELECT





Elastomer Manufacturing Process

Verification Process

• 100% camera visual inspection for pre-defined defects



End-of-Line Defect Reduction





100% Camera Inspection of rubber components

> CONNECTING PEOPLE SCIENCE AND REGULATION®

54 COPYRIGHT © PDA

technical report available – TR 2016-172



Automated vision inspection verification: defects examples





55 COPYRIGHT © PDA

pda.org



Automated vision inspection verification: defects examples







Elastomer Manufacturing Process

Steam sterilization

- Plungers, stoppers and lined seals
- The sterilization process is validated to assure a minimum SAL of 10⁻⁶ and in line with
 - ISO 17665-1 and 17665-2
- Steam processed elastomer formulations exhibit less degradation



Gamma sterilization

- > Plungers
- The sterilization process is validated to assure a minimum SAL of 10⁻⁶ and in line with
 - ISO 11137-1 and ISO 11137-2
- Gamma processing might impact degradation of the elastomeric formulation





Sterility assurance is reported in the quality certificate coming with every batch







Elastomer Manufacturing Process

Pack

• Product is packaged





Ship

• Prepare for final shipment to the customer







Differentiated Solutions: Increasing Quality & Inspection





pda.org



Particulates and Lack of Sterility Cause Most Product Recalls



attributable to container closure.



60 COPYRIGHT © PDA

¹ https://www.fda.gov/arugs/drug-safety-and-availability/drug-recalls (Accessed September 1, 2022) and https://www.fda.gov/vaccines-blood-biologics/safety-availability-biologics/recalls-biologics (Accessed October 17, 2022)





Visible and Subvisible Particle Specification Example

Attribute	NovaPure [®] 4023/50 Components			
Particulate > 5 μm < 10 μm	<100.0 partio	cles / 10 cm ²		
Particulate > 10 μm < 25 μm	<60.0 particles / 10 cm ²			
Particulate > 25 μm < 50 μm	<8.0 particles / 10 cm ²			
0.9 Particulate > 50 μm < 100 μm	<1.0 particl	es / 10 cm ²		
Particulate > 100 μm	<0.2 particles / 10 cm ²			
Fibers > 10 mm	AQL - 0.010	PPM ≤ 10		
Embedded Foreign Matter > 0.2 mm ²	AQL - 0.015	PPM ≤ 50		
Fibers ≥ 2 mm		PPM ≤ 250		
Defects potentially leading to non-sterility	AQL - 0.040	PPM ≤ 250		
Defects impairing processing		PPM ≤ 250		
Fibers ≥ 0.5 mm, < 2.0 mm	AQL - 0.25	PPM ≤ 2500		

Acceptable Quality Level (AQL) is a statistical measurement of the maximum acceptable number of defective goods in a particular sample size







Manufacturing Process



62 COPYRIGHT © PDA

pda.org



pda.org

Secondary Packaging







Secondary Packaging - Flexibility for Filling Needs

Filled bags are offered in ready-to-use (RU) quality by either steam or gamma validated processes

The ported bag packaging system is qualified to maintain the package integrity and stability of the components throughout the recommended shelf-life period. Verification includes shipping distribution simulation studies.















High-quality packaging materials

Reduction of particle load of primary packaging –	≻
tighter specification	

Ease of use

Pinhole resistant – physical – stress



Qualified to maintain the package integrity and stability of the components throughout the recommended shelf-life period. Verification includes shipping distribution simulation studies







Ready-to-Use Packaging Solutions







Relevant Compendial Chapters and Standards





pda.org

67^{OPYRIGHT}© PDA



68 COPYRIGHT © PDA

Global Comparison of Elastomer Chapters

Purpose	Paragraph	USP <381>	Ph Eur 3.2.9	JP 7.03	YBB
Introduction	Definition of Elastomer Types	\checkmark	\checkmark	-	\checkmark
Identification	e.g. IR, ash test	\checkmark	\checkmark	\checkmark	\checkmark
Physico-chemical Tests	Appearance of solution, absorbance, etc	\checkmark	\checkmark	\checkmark	\checkmark
Potential Extractable	Ammonium, Volatile Sulfides	\checkmark	\checkmark	\checkmark	\checkmark
Functionality Tests*	Fragmentation, self- sealing, …	\checkmark	\checkmark	-	\checkmark



pda.org



Global comparison of elastomer chapters









CONNECTING PEOPLE SCIENCE AND REGULATION®



Introduction to USP <1382> and <382>

<1382> Assessment of Elastomeric Component Functional Suitability in Parenteral Product Packaging/Delivery Systems

- Assist in the functional suitability assessment of elastomeric components as part of packaging / delivery systems
- ISO references
- Sampling plan guidance

<382> Elastomeric Component Functional Suitability in Parenteral Product Packaging/Delivery Systems

Fitness for intended use functional suitability tests and requirements

Released December 2020 with 5-year implementation grace period





Current <381> versus <382>

From: USP <381>

Elastomeric Closures for Injections

- Functionality Tests
 - Penetrability
 - Fragmentation
 - Self-Sealing Capacity

Container Closures for Vials and Bottles



To: USP <382>

Elastomeric Component Functional Suitability in Parenteral Product Packaging and Delivery Systems

- Package/Delivery System Integrity Tests
- Needle and Spike Access Functionality Tests
 - Fragmentation
 - Penetration Force
 - Needle Self-Sealing Capacity
 - Spike Retention and Sealability Capacity
- Plunger Functional Suitability Tests
 - Plunger Break Force and Plunger Glide Force
 - Plunger Seal Integrity
- Tip Cap and Needle Shield Functionality Tests

System Closures for Vials, Bottles, Blow Fill Seal Containers, Plastics, Cartridges and Syringes




Thank you very much for your attention!

Any Thoughts? Any Questions?

Christa.Jansen-Otten@westpharma.com

West and the diamond logo, NovaPure, VeriSure, Sterilizablebag, FluroTec, Envision, and Westar are trademarks or registered trademarks of West Pharmaceutical Services, Inc. in the United States and other jurisdictions. Biosafe is a trademark of Sartorius Stedim Aseptics. DPTE BetaBag is a trademark of GETINGE LA CALHENE. All other trademarks appearing in this presentation are the properties of their respective owners. FluroTec and B2-Coating technologies are licensed from Daikyo Seiko, Ltd.



73 COPYRIGHT © PDA