

GMP for Pre-Filled Syringes (PFS)

Development, Manufacturing, Control
Part of PharmaCongress 2024

19/20 March 2024

Wiesbaden, Germany

With a view on
the implications
of the New EU
GMP Annex 1!

Highlights

- Basics & Regulatory Overview
- Pre-fillable Syringes Design and Requirements
- Fill-Finish & Assembly Processes
- Process Simulation / Validation
- Visual Inspection & Container Closure Integrity
- Contamination Control Strategy
- MDR - Ensuring Compliance for Syringe based Combination Products
- GMP issues in Inspections
- Case Studies

Speakers

Maria Luisa Bernuzzi | MesaLabs, France

Jean-François Decoster | UCB, Belgium

Katharina Golly | Novartis, Switzerland

Dr Bernhard Illes | Microcoat Biotechnologie, Germany

Christa Jansen-Otten | West, Germany

Dr Daniel Müller | Local GMP Authority of Baden Württemberg, Germany

Dr Wenzel Novak | Gerresheimer Bünde, Germany

Jinesh Sadalge | Novartis, Austria

Dr Helen Sauter | Vetter Pharma-Fertigung, Germany

Dr Max Scheible | Vetter Pharma-Fertigung, Germany



This conference is part of PharmaCongress 2024



Academy
Your GMP/GDP
Information Source



OBJECTIVES

In this conference you will learn which requirements for pre-fillable syringes are defined by the regulations. You get to know all aspects of the manufacture of pre-fillable syringes that influence the filling process and the quality of the final product. In addition, practice-oriented case studies will guide you through the relevant production processes, simulations and controls for pre-filled syringes.

BACKGROUND

Currently there is a growing demand in the development of pre-fillable syringes (e.g. ready-to-fill, ready-to-use, sterile clean filling) for several enhanced Biotech applications (i.e. for the final product, the Pre-filled Syringe). However, new GMP requirements,

also for the sterile packaging material (e.g. regarding validation of the sterilization procedure for the syringe), apply with the revised EU GMP Annex 1 entitled "Manufacture of Sterile Medicinal Products".

This event will therefore deal with the current discussions and trends in the manufacture of pre-filled syringes:

- GMP requirements for pre-fillable syringes / devices
- PFS Design & Safety Systems
- Alternatives to glass
- GMP Requirements for personnel, cleanrooms, equipment & facilities
- Processing of pre-filled syringes
- Auto-injector Assembling

PROGRAMME 19 March 2024

Regulatory Overview, Annex 1 Impact and Inspection experience

Dr Daniel Müller, *Local GMP Authority of Baden-Württemberg, Germany*

- Regulatory framework (EU), impact for pre-filled syringes
- Impact of new Annex 1
- Inspection experience

Contamination Control Strategy

Dr Helen Sauter, *Vetter Pharma-Fertigung*

Practical experiences

- CCS – a new Annex 1 requirement
- Case Study: CCS implementation
- Risk based approach for control point identification

Medical Device Regulations - Understanding the Impacts and ensuring Compliance for Syringe-based Combination Products

Christa Jansen-Otten, *West*

- Navigating the EU MDR Regulations requirements
- Advantages of platforming on prefillable syringes
- Case example of technology being applied by the market for platform applications
- Needs of suppliers for supportive documentation

PFS made from Glass or Polymer

Katharina Golly, *Novartis*

- Materials
- Manufacturing
- Sterilization methods
- Design
- Pros and Cons

PFS and Needle Safety Systems

Katharina Golly, *Novartis*

Jinesh Sadalge, *Novartis*

- Regulatory Requirements
- Active vs. Passive Systems
- Design Considerations
- Examples

Validation of a Steam Sterilization Process for a Pre-Filled Syringe

Maria Luisa Bernuzzi, *MesaLabs*

- Challenges in steam sterilization of a PFS and its biological validation
- How to manage a heat sensitive load
- Bioburden/biological indicators approach, D value determination and the correct choice of biological indicators
- Validating the specific cycle



- Contamination Control Strategy
- Observations during GMP inspections

The presentations will be provided in a practice-oriented way from the different viewpoints of authorities, suppliers of packaging materials / devices / services (including sterilization activities), and the pharmaceutical industry.

TARGET AUDIENCE

This event is designed for employees working in the pharmaceutical industry (including drug device combination products), for suppliers of packaging materials & devices and to all who have to deal with the manufacture, control and release of prefilled syringes.

They key areas are

- Sterile Production
- Packaging material / Device development
- Manufacturing / Processing / Assembling
- Quality Control
- Quality Assurance

MODERATOR

Dr Wenzel Novak, *Gerresheimer Bünde*

PROGRAMME 20 March 2024

Container Closure Integrity

Jean-François Decoster, *UCB*

- Requirements for CCIT
- Method development and validation

Process Simulation / Media Fill

Dr Helen Sauter, *Vetter Pharma-Fertigung*

- Media Fill Design
- Worst-case parameters & requirements
- Validation of processes with Media Fills
- Trends with regards to Media Fills

Visual Inspection

Jean-François Decoster, *UCB*

- Requirements
- Method development and validation
- AQL testing
- Automated vs. semi-automated vs. manual inspection

Automated Visual Inspection: Process and Transfer

Dr Max Scheible, *Vetter Pharma-Fertigung*

- Automated Visual Inspection (AVI) as an alternative to MVI
- State-of-the-art technologies for a robust and reproducible process
- Qualification & Transfer

Endotoxin Detection in Pre-Filled Syringes: Challenges during Method Development and Validation

Dr Bernhard Illes, *Microcoat Biotechnologie*

- Introduction to Endotoxin testing and endotoxin masking (Low Endotoxin Recovery (LER))
- General approach for development and validation of endotoxin detection methods
- Considerations and challenges for method development and validation for PFS
- Case studies for method development and validation for GMP release testing

SPEAKERS



Maria Luisa Bernuzzi

MesaLabs, France

Maria is working as Product and Application Engineer for MesaLabs. She has deep knowledge and experience in validations of steam, dry heat sterilization/ depyrogenation and hydrogen peroxide decontamination to guarantee the microbiological quality of the product.



Jean-François Decoster

UCB, Belgium

Jean-François holds a Master Degree in Chemical Engineering from the Brussels Industrial Superior School. He joined UCB in 2005 where he took increasing responsibilities in Primary Packaging Development. Since 2010, he has been the Head of Primary Packaging Development for UCB. In 2022, he moved to the Global Quality organization of UCB, where he is now in charge of several strategic projects, including Annex 1 implementation.



Katharina Golly

Novartis, Switzerland

Katharina is Senior Expert Engineering and began her professional career at Schott. Among other things, she was responsible for the development of silicone-based coatings for prefilled glass syringes. In 2015, she moved to Novartis as a packaging expert and supported ophthalmic PFS projects before becoming the technical lead for vials & kits.



Dr Bernhard Illes

Microcoat Biotechnologie GmbH, Germany

Bernhard joined Microcoat as a project manager in 2021 and is in charge of endotoxin service projects with a focus on LER, mitigation, method development and validation.



Christa Jansen-Otten

West, Germany

Christa is currently Director of Technical Product Development at West Pharmaceutical Services Inc. She has worked within the pharmaceutical industry for more than 20 years and gained experience as QA Manager in one of the world's leading pharmaceutical companies in sterile filling and packaging.



Dr Daniel Müller

GMP/GDP Inspector, Local Government, Germany

Daniel is currently head of the GMP Inspectorate at the local competent authority (GMP inspectorate) in Tuebingen, Germany. Since 2001 he has been working as GMDP inspector, conducting national inspections as well as EMA and overseas inspections. Before joining the authority, Dr Müller was working in the pharmaceutical industry, last serving as Qualified Person for sterile drug products.



Dr Wenzel Novak

Gerresheimer Bünde, Germany

Wenzel studied biology and did his PhD in physics at the Max-Planck-Institute for Neurochemistry in Munich. As head of laboratory in a Swiss biotech-company, he worked on Keratinocytes for wound healing. In charge of project management and as head of production for pre-sterilized syringes, he designed the manufacturing area, built up the process and quality systems and operated the start-up phase of production. He took role as Chief-Innovation-Officer at a pharmaceutical equipment supplier, developing new methods of sterilization and filling processes. Two years in USA, he developed the market for cell- and gene therapy equipment. In 2018, he returned to the same primary packaging manufacturer, left 13 years ago in a new global senior role for business development.



Jinesh Sadalge

Novartis, Austria

Jinesh is a Senior Expert Engineering and started career as a Validation Engineer at Barry Plastics (Formerly known as REXAM) in 2012. There he was responsible for the validation of injection molding, assembly lines and test methods. Before joining Novartis, he worked with Biocon and Medtronic as Design Quality Engineering for Pen injectors, Auto injectors and Spinal cord stimulators. In 2020, he changed as a Device Manager to Novartis and working on Needle Safety Devices projects before taking the Delivery System lead for Syringe and Safety system in 2023.



Dr Helen Sauter

Vetter Pharma-Fertigung, Germany

Helen received her Ph. D. in microbiology at the University of Stuttgart-Hohenheim. She has been working for Vetter since 2013. Currently she holds the position of Director QA – Sterility Assurance/Lab Operation/Training systems.



Dr Max Scheible

Vetter Pharma-Fertigung, Germany

Max made his PhD in physics at the Technical University Munich in 2014, specializing on DNA nanotechnology. Afterwards he worked as a post-doc at the Technical University Braunschweig and co-founded an EXIST-based company in the field of microscopy. In 2018 he joined Vetter as part of the Development Service. Since then he is responsible for process development and process qualification as well as the implementation of new AVI products.

The guiding theme of the GMP PharmaCongress 2024 on 19/20 March will be „users sharing challenges and solutions in practice“. Therefore, benefit from your colleagues’ experience and from the direct information exchange at the GMP PharmaCongress & GMP PharmaTechnica 2024.

The Conference Tracks

As a participant you can switch between any of the **11 conference tracks** any time and also visit the PharmaTechnica Expo with more than 110 international exhibitors.

The GMP PharmaCongress Conference Tracks	19 March 2024	20 March 2024
Non-Sterile Products – Challenges in Manufacturing & Quality	✓	n.a.
GMP – Green or Good Manufacturing Practice?	✓	n.a.
Packaging/Packaging Materials – Challenges & Solutions	n.a.	✓
European Aseptic Conference – Technology	✓	✓
Trends in Barrier Systems & Robotics	✓	✓
Modern Cleanroom Technology	✓	✓
Digitalisation & Artificial Intelligence	✓	✓
GMP for Pre-Filled Syringes (PFS)	✓	✓
Lyophilization – Modern Techniques & New Requirements	✓	✓
ATMPs – Hurdles & Achievements in Quality and Safety	✓	✓
Vaccines – Advantages & Challenges in Manufacturing	✓	✓
GMP PharmaTechnica Expo	✓	✓

Keynote on 19 March 2024

Manufacturing of Pandemic Vaccines – Manufacturing & Supply Solutions Enabling the Delivery of Large Numbers of Vaccine Doses

Dr Guido Dietrich, CEPI

Keynote on 20 March 2024

Presentation by the Wallhäußer Innovation Award Winner

In the Live Demo Area in the PharmaTechnica Expo hall you will benefit from the exhibitors’ demonstrations – presenting their latest technology, products and services. Take advantage of these live performances – and get to feel and experience their products. For a list of all companies exhibiting at PharmaTechnica, please see the exhibitor list and plan on the website at www.pharma-congress.com.

Day	Time	Exhibitor	Stand	Live Demo
19 March 2024	10.00-10.15 h	Ellab	A 16	Self mapping made easy
	10.15-10.30 h	boTec	A 7	Optimized planning and operation of pharmaceutical storage and distribution systems
	11.15-11.30 h	Bausch+Ströbel	A 33	OMNIA - Boost your processes and step closer to your pharmaceutical plant of the future
	11.30-11.45 h	Merck	B 32	Annex 1's "Specific Risks Associated with Single-Use Systems"
	11.45-12.00 h	Cytiva	C 5	Point-of-use leak testing of single-use systems
	12.30-12.45 h	MK Versuchsanlagen	A 12	State of the art testing of barrier systems
	12.45-13.00 h	Innerspace	B 14	Aseptic Training with Virtual Reality
	13.00-13.15 h	MBV	B 20	MBV MAS-100 ISO – Microbial Air Sampler
	15.00-15.15 h	ZETA	B 12	Intelligent solution for passing single-use tubes through cleanroom walls
	15.15-15.30 h	Friedrich Sailer	B 31	Shining Solutions: Live Demo of Cutting-Edge Stainless Steel Cleanroom Equipment
15.30-15.45 h	Particle Measuring Systems	C 7	Facility Monitoring Systems	
20 March 2024	09.45-10.00 h	PHARMAPLAN	A 37	Virtual Pharma Campus
	10.00-10.15 h	Emerson Automation Solutions	B 22	Real-Time Scheduling and Production Optimization
	10.15-10.30 h	Yokogawa	C 9	Predict, Prevent, Perform: A Proactive Approach to Asset Health in the Pharmaceutical Industry
	12.00-12.15 h	REA Elektronik	B 29	Experts in Printing and Code Verification of pharmaceutical and medical-device packaging (f.e.UDI/MDR)
	12.15-12.30 h	IWT / Tecniplast	B 21	High pressure cleaning in pharmaceutical production. Advantages, challenges, sustainability and savings
	12.30-12.45 h	Quascenta Pte	B 6	Using an Intelligent Lifecycle-Based Solution for Streamlined Product Tracking, Design Optimization, Collaboration, and Reporting
	12.45-13.00 h	Kneat Solutions	C 12	Kneat Gx Demo

Live Demos and Times Subject to Change

Easy Registration

Registration Form:
CONCEPT HEIDELBERG
Rischerstraße 8
69123 Heidelberg

Registration Form:
(06221) 84 44 34

E-Mail:
info@concept-heidelberg.de

Internet:
www.pharma-congress.com

Congress Dates

Tuesday, 19 March 2024, 09.00 - 18.00 h
Wednesday 20 March 2024, 09.00 - 17.00 h
Registration

Tuesday & Wednesday, 19/20 March 2024, 08.00 - 09.00 h

Fees

The one day ticket is available for € 690,- plus VAT, both days for € 1,380.- plus VAT. It includes participation in any track on that day and the visit of the PharmaTechnica Expo. In addition, lunch and beverages during the tracks and in breaks as well as the social event on the evening of the first congress day, 19 March is included; please mark if you would like to attend the Social Event. The fee is payable in advance after receipt of invoice.

Venue

RheinMain CongressCenter (rmcc)
Friedrich-Ebert-Allee 1 | 65189 Wiesbaden
Phone: +49 (0) 611 1729-444
E-Mail: veranstaltungsservice-rmcc@wicm.de

Accommodation

CONCEPT HEIDELBERG has reserved a limited number of rooms close to the CongressCenter. You will receive a room reservation form when you have registered for the conference. Reservation should be made directly with the hotel. Early reservation is recommended.

Registration

Via the attached reservation form, by e-mail or by fax message. Or you register online at www.gmp-compliance.org.

Presentations/Certificate

The presentations for this event will be available for you to download and print before and after the event. Please note that no printed materials will be handed out on site and that there will not be any opportunity to print the presentations on site. After the event, you will automatically receive your certificate of participation.

Conference language

The official conference language will be English.

Organisation and Contact

ECA has entrusted Concept Heidelberg with the organisation of this event.
CONCEPT HEIDELBERG
P.O.Box 10 17 64
69007 Heidelberg, Germany
Phone: +49 (0) 62 21 / 84 44-0 | Fax: +49 (0) 62 21 / 84 44 34
info@concept-heidelberg.de | www.concept-heidelberg.de

For questions regarding content please contact:
Dr Andrea Kühn-Hebecker (Operations Director) at
+49 (0) 62 21 / 84 44 35, or at kuehn@concept-heidelberg.de

For questions regarding organisation please contact:
Mr Ronny Strohwald (Organisation Manager) at
+49 (0) 62 21 / 84 44 51, or at strohwald@concept-heidelberg.de

If the bill-to-address deviates from the specifications on the right, please fill out here:

CONCEPT HEIDELBERG
P.O. Box 101764
Fax +49 (0) 62 21/84 44 34
D-69007 Heidelberg
GERMANY

Reservation Form (Please complete in full)

GMP for Pre-Filled Syringes (PFS)

Part of PharmaCongress 2024

19/20 March 2024, Wiesbaden, Germany

- Day 1 & 2 (19/20 March 2024)
 Day 1 (19 March 2024)
 Day 2 (20 March 2024)
 Yes, I would also like to take part in the Social Event on the evening of 19 March 2024.

Mr Ms Mx Dr

First name, Surname

Company

Department

Important: Please indicate your company's VAT ID Number

P.O. Number (if applicable)

Street/P.O. Box

City

Zip Code

Country

Phone/Fax

E-Mail (please fill in)

General terms and conditions

If you cannot attend the conference you have two options:

1. We are happy to welcome a substitute colleague at any time.

2. If you have to cancel entirely we must charge the following processing fees:

- Cancellation until 4 weeks prior to the conference 10 %

- Cancellation until 3 weeks prior to the conference 25 %

- Cancellation until 2 weeks prior to the conference 50 %

- Cancellation within 2 weeks prior to the conference 100 %

CONCEPT HEIDELBERG reserves the right to change the materials, instructors, or speakers without notice or to cancel an event.

If the event must be cancelled, registrants will be notified as

soon as possible and will receive a full refund of fees paid.

CONCEPT HEIDELBERG will not be responsible for discount airfare penalties or other costs incurred due to a cancellation.

Terms of payment: Payable without deductions within 10 days after receipt of invoice.

Important: This is a binding registration and above fees are due in case of cancellation or non-appearance. If you cannot take part, you have to inform us in writing. The cancellation fee will then be calculated according to the point of time at which we receive your message.

In case you do not appear at the event without having informed us, you will have to pay the full registration fee, even if you have not made the payment yet. Only after we have received

your payment, you are entitled to participate in the conference

(receipt of payment will not be confirmed)! (As of July 2022).

German law shall apply. Court of jurisdiction is Heidelberg.

Privacy Policy: By registering for this event, I accept the processing of my Personal Data. Concept Heidelberg will use my data for the processing of this order, for which I hereby declare to agree that my personal data is stored and processed. Concept Heidelberg will only send me information in relation with this order or similar ones.

My personal data will not be disclosed to third parties (see also the privacy policy at http://www.gmp-compliance.org/eca_privacy.html). I note that I can ask for the modification, correction or deletion of my data at any time via the contact form on this website.