



PDA Manufacturing Meets Legislation Workshop 2024  
Berlin, Germany  
04-06 September 2024  
Change Hub, Hardenbergstraße 32, 10623 Berlin

# Agenda

Wednesday, 04 September 2024

13:00 - 17:00 CEST

Module 1: Preparation for EMA Inspectors Working Group Dialogue		Moderator: <i>RAQAB Representative</i>
13:00	Welcome	Falk Klar, <i>PDA</i>
	Introduction to Industry-Regulators Interaction	<i>RAQAB Representative</i>
	Presentation of the Submitted Topics Workshop Speed Discussions	<i>All Topic submitters</i>
15:15	Coffee Break	



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	Discussion Round with Readout	<i>All Topic Submitters</i>
	Summary of Selected Topics	<b>Moderator:</b> RAQAB Representative
17:00	End of Workshop Day 1	
<b>Thursday, 05 September 2024</b>		<b>09:00 - 16:45 CEST</b>
<b>Module 2: Impact of Planned PFAS Ban on the Pharmaceutical Industry in Europe</b>		<b>Moderator:</b> Ana Kuschel, <i>WEST</i> Ankur Kulshrestha, <i>BMS</i>
09:00	PFAS Materials Used in Pharmaceutical Manufacturing Settings – Potential Supply Chain Risk	Louise Loughran, <i>Eli Lilly and Company</i>
	Overview of the PFAS Situation for their Use in Pharmaceutical Filtration, Opportunities and Risks	Alexander Mitropoulos, <i>Merck</i>
	Considerations for Early-Stage Testing of Chemicals for PFAS Replacement	Dieter Hennecke, <i>Fraunhofer Institut</i>
10:30	<b>Coffee Break</b>	
	Flipchart Activity on PFAS Risk Assessment	<i>Invited Speakers</i>
12:00	<b>Lunch Break</b>	



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<b>Module 3: Challenges and Opportunities in Pharmaceutical Manufacturing in Europe</b>		<b>Moderator:</b> Veronique Davoust, <i>Pfizer</i>
13:00	EDQM Remote Inspections	Oisín Daly, <i>EDQM</i>
	Challenges of a Public Affairs Perspective – Values, Mission, and Patient Centricity	<i>Merck Public Affairs</i>
	Innovative Manufacturing Approaches in Europe: From Idea to Implementation	Ursula Busse, <i>Tigen Pharma</i>
	PDA’s Initiative for Early Career Professionals	Bettine Boltres, <i>WEST</i>
	Challenges and Opportunities	<i>Invited Speaker</i>
<b>14:45</b>	<b>Coffee Break</b>	
	Road to Resilience: A Young Professional's Journey	Miriam Kremer van der Kamp, <i>Vils</i>
	Running the Pharmaceutical Packaging Race	Ana Kuschel, <i>WEST</i>
	<b>Panel Discussion</b>	
<b>16:45</b>	<b>End of Workshop Day 2</b>	



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Friday, 06 September 2024		09:00 - 11:15 CEST
<b>Module 4: Overview of the Current Changes in the EU Regulatory Framework</b>  <i>This session will provide an overview of the hot topics changing in the EU legislative framework impacting manufacturing, quality, and supply.</i>		<b>Moderator:</b> Stephan Rönninger, Amgen
<b>09:00</b>	<b>Welcome Coffee Break</b>	
	Pharmaceutical Manufacturing in the Regulatory Framework	Stephan Rönninger, Amgen
	Overview of Upcoming Regulatory Changes Relating to Sustainability	Marion Briggs, Airfield Consulting
	Getting to Pharma Organizational Action on Sustainability	Joachim Almdal, Green Innovation Group
<b>Panel Discussion</b>  <b>General Pharma Legislation</b>  <i>Impact on CMC, manufacturing, environmental risk assessment, inspections, imported products, and supply.</i>  <b>‘License to Operate’</b>  <i>Under that term, the requirements for environmental control and sustainability are managed</i>  <b>Drug Shortage Prevention</b>  <i>While companies and marketing authorization holders follow basic principles implementation measures by individual EU Member States and other countries may benefit from a holistic harmonized approach and identifying the real root causes.</i>		<b>Moderator:</b> Stephan Rönninger, Amgen



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	Key Messages: Call to Action	<b>Moderator:</b> Stephan Rönninger, <i>Amgen</i>
	Farewell	Falk Klar, <i>PDA</i>
<b>11:15</b>	<b>End of Workshop</b>	

The agenda is subject to change.