

# ExpertFORUM Labelling

Attendance-based and online: Essential trends in product information, digitalisation, safety labelling & IDMP

## TOPICS

- Implementation of IDMP – current status
- QRD Working Group update
- Electronic product information (ePI) and Germany's GI 4.0
- E-labelling projects and digitalisation
- Labelling compliance and pharmacovigilance
- Accessibility of product information

## YOUR SPEAKERS

### **Dr Peter Bachmann**

Senior Expert Regulatory Affairs, GERMANY

### **Berit Fasse, MDRA**

Bayer AG, GERMANY

### **Dr Eva Göbgen**

vfa Association of Research-Based Pharmaceutical Companies, GERMANY

### **Dr Thomas Grüger**

Senior Expert Pharmacovigilance, GERMANY

### **Dr Niklas Jänich**

Boehringer Ingelheim International GmbH, GERMANY

### **Karl-Heinz Loebel**

PharmaLex GmbH, GERMANY

### **Kim Sherwood**

Medical Products Agency (MPA), SWEDEN

### **Hjörleifur Thorarinsson**

Icelandic Medicine Verification Organisation, ICELAND

### **Dr Anke Webler-Messenger**

Boehringer Ingelheim International GmbH, GERMANY

## **YOUR SPEAKERS DAY 1**



**Dr Peter Bachmann**

Bonn, GERMANY

Senior Expert Regulatory Affairs



**Dr Eva Göbgen**

German Association of Research-based Pharmaceutical Companies (vfa e.V.),  
Berlin, GERMANY

She manages the project GI 4.0, the German pilot for digital patient leaflets.



**Dr Niklas Jänich**

Boehringer Ingelheim  
International GmbH, Ingelheim,  
GERMANY

Head of Global Reg. Affairs Processes/  
Training/Compliance & Global Labeling  
Operations



**Kim Sherwood**

Medical Products Agency (MPA),  
Uppsala, SWEDEN

Senior Expert Product Information Assessor,  
Department of Product Information



**Hjörleifur Thorarinsson**

Icelandic Medicine Verification  
Organisation,  
Reykjavík, ICELAND

Executive Director

## **PROGRAMME DAY 1**

### **Implementation of IDMP – current status**

Karl-Heinz Loebel

- The development of IDMP and SPOR

### **QRD Working Group update**

Kim Sherwood

- ePI and QRD templates – fragments, structures and XML
- Traceability of biologics
- Revision of the SmPC guidelines

### **Electronic product information (ePI) – current status in the EU**

Dr Peter Bachmann

### **E-labelling projects and digitalisation – Icelandic Medicines Verification Organisation (ICEMVO) update**

Hjörleifur Thorarinsson

### **GI 4.0 – the German project**

Dr Eva Göbgen

- Concept of the German ePI pilot project
- Source of information
- What happens next
- ‘Rote Liste’ in Germany

### **Labelling content management**

Dr Niklas Jänich

- Overcoming the document paradigm
- Leveraging structured content management for the E2E labelling process
- Structured authoring tool development
- IDMP compatibility

## **PROGRAMME DAY 2**

### **Labelling compliance and pharmacovigilance**

Dr Thomas Grüger

- Regulatory duty to update the package leaflet after signal detection following the PRAC decision
- Educational material as an authority requirement
- Variations submission and implementation
- Labelling as a trigger or part of PV inspections

### **Assuring labelling compliance in practice**

Dr Anke Webler-Messenger

- Continuous updating of company core safety data and company core safety information, possible triggers and challenges
- Dealing with and avoiding local labelling deviations
- Roles, responsibilities and mechanisms to assure labelling compliance – governance bodies and decision-making
- Demonstrating labelling compliance during PV inspections

### **Patient communication beyond readability testing**

Berit Fasse

### **Collecting, cleaning and structuring data in companies**

Karl-Heinz Loebel

- ... for IDMP
- ... to improve labelling

### **PDF/UA: accessibility for people with disabilities using assistive technology**

Karl-Heinz Loebel

## **YOUR SPEAKERS DAY 2**



**Berit Fasse**

Bayer AG, Wuppertal,  
GERMANY

Senior EU Labeling Manager



**Dr Thomas Grüger**

Bonn, Germany

Senior Expert Pharmacovigilance



**Karl-Heinz Loebel**

PharmaLex GmbH,  
Mannheim, GERMANY

Director, Principle Consultant Regulatory Operations



**Dr Anke**

**Webler-Messenger**

Boehringer Ingelheim  
International GmbH,  
Ingelheim, GERMANY

Head of Regulatory Affairs CNS,  
Retinopathy and Emerging Areas

## AIMS AND OBJECTIVES

This ExpertFORUM will provide a thorough update on current trends in product information and labelling. Authority and industry experts will talk about e-labelling, digitalisation and the identification of medicinal products (IDMP), and give you practical examples of labelling content management and compliance.

A discussion on the accessibility of product information to patients, including tips on how to optimise your labelling management, will round off the event.

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## TIME SCHEDULE

Tuesday, 8 December 2020  
from 08:30 registration  
from 09:00 – 17:00 conference

Wednesday, 9 December 2020  
from 09:00 – 16:00 conference

## WHO SHOULD ATTEND

This conference addresses the needs of individuals involved in product information in regulatory affairs, medical affairs and pharmacovigilance.

It will provide useful updates on the relevant trends in digitalisation, compliance and much more. A working knowledge of product information is a prerequisite.

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## YOUR SAFETY AND THE OPTION TO PARTICIPATE ONLINE

For the first time, participation in the ExpertFORUM Labelling conference will be both attendance-based and online. On site, we will ensure that the necessary safety precautions are met and use significantly larger rooms and the latest hygiene measures. At the same time, a professional streaming service provider is on board to make the conference accessible from your own workplace. Direct interaction with the speakers in the conference room will also be possible.

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## REGISTRATION FORM

**service@forum-institut.de**

**www.forum-institut.de**

**Webcode 2012234**

**Tel. +49 6221 500-500**

**Fax +49 6221 500-555**

### Venue

Heidelberg Marriott Hotel  
Tel. +49 6221 908-0

or in your office / homeoffice

### Fee

1,890.00 (+German VAT)



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## CANCELLATION POLICY

Our general terms and conditions apply (as of 1. January 2016) and are available upon request. We can send them to you anytime or you can find them on the internet at [www.forum-institut.de/agb\\_en](http://www.forum-institut.de/agb_en)

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## ANY FURTHER QUESTIONS?



### Dr. Henriette Wolf-Klein

Bereichsleitung  
Pharma & Healthcare  
Tel. +49 6221 500-680  
[h.wolf-klein@forum-institut.de](mailto:h.wolf-klein@forum-institut.de)