

Masterclass EU QPPV

Current challenges for the Qualified Person for PV

Topics

- Efficient delegation and cooperation with your deputy
- How to maintain oversight of multinational PV activities

- Your key to success with many practical tips for your daily work.
- Sustainable collaboration with other departments
- Mergers and acquisitions: how do you ensure that PV-compliance is not compromised?
- Workshop: Resource planning in pharmacovigilance

Your speakers



Dr Katharina Caspary Horizon Pharma GmbH, Mannheim



Dr Monica Rusu Abbott Laboratories GmbH, Hannover

Aims and objectives

The duties of the European Qualified Person for Pharmacovigilance are diverse and challenging. This is why our advanced seminar will provide you with practical tips and hints for your work, and further equip your available toolbox.

After the seminar, you will be able to:

- set up an efficient QPPV delegation and deputy system;
- work more effectively together with other important departments to ensure the quality of your PV system;
- maintain oversight of multinational PV acitivities; and
- plan the resources you need in a changing regulatory environment.

Who should attend?

This seminar is intended for those working in the pharmaceutical industry that have an extensive regulatory background in terms of the QPPV role and now would like to improve their daily processes.

The following will especially benefit from this seminar:

- EU QPPVs and their deputies;
- National Persons Responsible for Pharmacovigilance;
- Heads of Pharmacovigilance; and
- consultants providing QPPV services.

Your speakers



Dr Katharina Caspary Horizon Pharma GmbH, Mannheim

Director Pharmacovigilance;
Medical doctor with profound experience
in various areas in the pharmaceutical
industry. Proven track-record of successful
strategic roles in pharmacovigilance and
clinical research and well-experienced
in cross-functional projects as well as
M&A across Medical Affairs, Regulatory,
Marketing & various outside parties.



Dr Monica Rusu Abbott Laboratories GmbH, Hannover

Director PV Governance & EU QPPV, Global Pharmacovigilance Innovation & Development, Established Pharmaceuticals

International courses

Are you interested in our international education programme? We provide a variety of specialised courses for the healthcare industry.

Quality guaranteed!

We follow the IMI quality criteria. An evaluation of participants' feedback on our healthcare courses produced a result of 1.7 (school grading system of 1-6).

Your programme

> 9.00 Welcome and Introduction round

> 9.15

National Person Responsible for PV vs EU QPPV

Dr Monica Rusu

- Regulatory background in Europe
- Responsibilities in SMEs and large companies
- Liability
- How to handle interfaces
- > 10.00 Coffee break

> 10.15

Workshop: Efficient delegation

Dr Monica Rusu

- Mandatory and obligatory tasks to delegate
- Correct documentation
- Cooperation with your deputy
- Establish your own monitoring system
- > 12.00 Lunch

> 13.00

How to maintain oversight of multinational PV activities

Dr Katharina Caspary

- PSMF the most important document?
- How to keep up-to-date with all the national peculiarities in Europe
- Useful processes and company structures
- The correct way to handle all the data and information
- Gathering all the information from license- and PV-partners
- The QPPV's role in ensuring affiliate compliance

> 14.30

Collaboration with other departments

Dr Monica Rusu

- Important contact points in the company: Quality, Clinical, Commercial, Medical Affairs, Regulatory Affairs best practices to ensure QPPV involvement
- Quality deficiencies: responsibility of the EU QPPV? How to organize possible recalls
- Harmonised quality system in PV, QA and Medical Affairs - a possible approach?
- > 15.30 Coffee break

> 15.45

Mergers and acquisitions: How do you ensure that PV-compliance is not compromised?

Dr Katharina Caspary

What has to be considered after an acquisition?

> 16.15

Workshop: Resource planning in Pharmacovigilance

Dr Katharina Caspary, Dr Monica Rusu

- What resources do you need for a working PV system
- Manpower, and workload
- > 17.00 End of seminar

Registration under service@forum-institut.com or Fax +49 6221 500-555

Registration Form

Yes, I will attend the seminar
☐ Masterclass EU QPPV

Name	
Position/Department	
Company	
Street	
Postal Code/City/Country	
Tel. No.	
E-Mail	
Contact person at office	
Date, Signature	

How to register

Registration: +49 6221 500-500

Conference-No. 17 09 203

Date/Venue:

Monday, 25 September 2017 in Frankfurt 8.30 registration; 9.00 - 17.00 seminar relexa hotel Lurgiallee 2 · 60439 Frankfurt Tel. +49 69 95778-0 · Fax +49 69 95778-876

Fee:

€ 990.00 (+ German VAT)

The fee includes course documentation (incl. free download) as well as mid-session refreshments, lunch and certificate. Invoice and confirmation will be forwarded to you.

Hotel accommodation:

A limited number of rooms have been reserved at the hotel and are subject to availability. Please book at least six weeks prior to the seminar to obtain a hotel room at the discounted rate. All bookings should be made directly with the hotel quoting FORUM Institut and the Course No.

Any Further Questions?



Please feel free to contact me if you have any questions.

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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.com/t&c