

## “Excellence in Pharmacovigilance” Module 3: Quality Management System

Module 3 of the Joint DIA/MEB “Excellence in Pharmacovigilance” covers establishment of a Quality Management System in Pharmacovigilance.

It includes aspects of the applicable GVP modules, as well as preparation and conduct of audits and inspections. This module also addresses risk-based assessments, document management and PV agreements.

### Day 1      17 Nov, 08:30-13:00

08:30 Welcome

09:00 Pharmacovigilance System Master File (PSMF)

*Pieter Grotenhuis, Senior Inspector for Pharmacovigilance, Health and Youth Care Inspectorate, Netherlands*

10:00 Audits and Inspections in Pharmacovigilance – Regulatory Perspective

*Pieter Grotenhuis, Senior Inspector for Pharmacovigilance, Health and Youth Care Inspectorate, Netherlands*

11:00 Break

11:30 Audits and Inspections in Pharmacovigilance – Industry Perspective

*Wendy Huisman, Director, Vigifit, Netherlands*

12:30 Discussion and Q&A

13:00 End of Day 1

### Day 2      18 Nov, 09:00-12:30

09:00 “Risk based” Activities, Documentation and Pharmacovigilance Agreements

*Wendy Huisman, Director, Vigifit, Netherlands*

10:30 Break

11:00 The Role of the Qualified Person Responsible for PV (QPPV)

*Wendy Huisman, Director, Vigifit, Netherlands*

12:00 Discussion and Q&A

12:30 End of Module 3

## “Excellence in Pharmacovigilance” Module 4: Risk Management

In accordance with the GVP Module V on Risk Management System, Risk Management Plans (RMPs) should be submitted by companies to propose activities aiming to identify, characterise or minimise risks associated with medicinal products. Given the potential public health implications and costs of such interventions, RMPs should be based on robust data. Examples will be discussed in this session.

Nonetheless, specific data collection and analysis are desirable for some safety issues, and pharmaco-epidemiological studies are the most performed tools of those “additional” Pharmacovigilance activities. This session also presents recent developments regarding risk communication.

### Day 1      22 Nov, 08:30-14:00

08:30 Welcome

09:00 Risk Management Plans – Regulatory Perspective: GVP Module V Revision 2

*Inge Zomerdijs, Pharmacovigilance Assessor, Medicines Evaluation Board, Netherlands*

10:00 Risk Management Plans – Industry Perspective

*Jan Petracek, Director, Institute of Pharmacovigilance, Czech Republic*

11:00 Break

11:30 Epidemiological Methods and Pharmacovigilance

*Fakhredin Sayed Tabatabaei, Senior Pharmacovigilance Assessor, Medicines Evaluation Board, Netherlands*

13:30 Discussion and Q&A

14:00 End of Day 1

### Day 2      23 Nov, 09:00-14:00

09:00 Effectiveness of Risk Minimisation Measures – Regulatory Perspective

*Inge Zomerdijs, Pharmacovigilance Assessor, Medicines Evaluation Board, Netherlands*

10:00 Effectiveness of Risk Minimisation Measures – Industry Perspective

*Jan Petracek, Director, Institute of Pharmacovigilance, Czech Republic*

11:00 Break

11:30 Harmonisation of RMP (HaRP) in Europe

*Paul ten Berg, Pharmacovigilance Assessor, Medicines Evaluation Board, Netherlands*

12:30 Risk Communication in EU – Challenges and Possibilities

*Jan Petracek, Director, Institute of Pharmacovigilance, Czech Republic*

13:30 Discussion and Q&A

14:00 End of Module 4

## “Excellence in Pharmacovigilance” Module 5: Signal Management

New safety signals may emerge at any time following product launch and must be evaluated for relative risk, medical importance, and likelihood of occurrence. Signal Management is, therefore, one of the crucial “routine” Pharmacovigilance activities. Approaches to Signal Management using qualitative and quantitative methods will be illustrated in a workshop with examples as well as general considerations on signal management in the EEA.

### Day 1      25 Nov, 08:30-13:00

08:30 Welcome

09:00 Introduction to Signal Detection in the European Union – Regulatory Perspective

*Bianca Mulder, Pharmacovigilance Assessor, Medicines Evaluation Board, Netherlands*

11:00 Break

11:30 Signal Management in the European Union – Industry Perspective

*Jan Petracek, Director, Institute of Pharmacovigilance, Czech Republic*

12:30 Discussion and Q&A

13:00 End of Day 1

### Day 2      26 Nov, 09:00-11:45

09:00 Signal Management – Workshop

*Bianca Mulder, Pharmacovigilance Assessor, Medicines Evaluation Board, Netherlands*

11:00 Discussion and Q&A

11:30 Closing Remarks

11:45 End of Module 5