EMA Excellence in Pharmacovigilance: Clinical trials and post-marketing training course

Course #14548
13-17 October 2014
European Medicines Agency, Churchill Place 30, Canary Wharf, London, UK

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Continuing Education
The Faculty of Pharmaceutical Medicine of the Royal Colleges of Physicians of the United Kingdom has accredited this training course with 25 CPD credits.
The Swiss Association of Pharmaceutical Professionals (SwAPP) and the Swiss Society for Pharmaceutical Medicine (SGPM) have accredited this training course with 34 credits.
This course has limited capacity. Register early

Overview
This course is designed to provide a firm grounding in key aspects of Global and mainly European Clinical Pre- and Post- Marketing Safety regulatory requirements. This five-day training course, presented by the European Medicines Agency, now also includes highlights and updates on the implementation of the new pharmacovigilance legislation and the latest news on the international harmonisation and standardisation activities in pharmacovigilance.

Who Will Attend
Professionals involved in pharmacovigilance and namely Qualified Persons for Pharmacovigilance (EU QPPV), clinical research, regulatory affairs, risk management, medical product safety assessment, and data analysis, epidemiology, labelling, quality assurance, compliance, medical information.

Level: Intermediate

Learning Objectives
For the five key topics as outlined below, the learning objectives also include the ability to:

• Describe the main changes to the business processes in the context of the new pharmacovigilance legislation
• Discuss the latest developments in the area of international harmonisation and standardisation with the main focus on the ICH E2B, E2C, E2F topics

At the conclusion of this course, participants should be able to:

Definitions and Methods in Pharmacovigilance
• Describe the scope and objectives of Pharmacovigilance and Risk Management and the relationship between the two concepts
• Discuss the development of definitions based on legislation and consensus fora
• Identify the key definitions and the vocabulary used in Pharmacovigilance in the European Union, illustrated by practical examples and exercises
• Explain and illustrate methods used in pharmacoepidemiology for measuring risks and estimating their association with drug exposure

Regulatory Aspects in Pharmacovigilance and Practical Examples
• Describe the European regulatory requirements in Pharmacovigilance
• Describe the requirements of establishing a Pharmacovigilance database and the use of the Medical Dictionary for Regulatory Activities (MedDRA) including the key functionalities of EudraVigilance
• Discuss the Pharmacovigilance System Master File and the preparation for audits and inspections

Diagnosis and Management of Adverse Drug Reactions
• Discuss the key elements of the medical evaluation of adverse events
• Recognise the important aspects in evaluating adverse events based on two examples
• Identify the main characteristics of two examples of drug induced adverse events

Signal Detection
• Understand MedDRA dictionary
• Describe signal detection and management in the EU, based on GVP module IX

Risk Management
• Explain the EU risk management strategy, the new approaches to risk assessment and prevention, and the different steps to be considered in the risk management process
• Describe the components of the Good Pharmacovigilance Practices (GVP) module V on the risk management systems
• Define the concept of risk, and explain differences between individual and population risks
• Describe current recommendations and practices of benefit-risk assessment, post-authorisation efficacy studies and post-authorisation safety studies
• Understand the main principles of risk communication based on case studies

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Monday | 13 October 2014

08:00 REGISTRATION

08:30 Keynote Presentation

THE ROLE OF THE EUROPEAN MEDICINES AGENCY IN PHARMACOVIGILANCE
Sabine Brosch, European Medicines Agency, EU

09:15 TOPIC 1

DEFINITIONS AND METHODS IN PHARMACOVIGILANCE
Topic 1 will provide a concise overview of the objectives and the scope of Pharmacovigilance and Risk Management and the relationship between the two concepts. The development of key definitions based on Community legislation and consensus fora such as the International Conference on Harmonisation of Technical Requirements for Registration of Pharmaceuticals for Human Use (ICH) and the CIOMS Working Groups will be summarised. Practical examples and exercises will be used to illustrate the key definitions and vocabulary applied in Pharmacovigilance.

09:15 Topic 1 Session 1

BASIC DEFINITIONS AND TOOLS IN PHARMACOVIGILANCE
Sabine Brosch, European Medicines Agency, EU
Gaby Danan, Pharmacovigilance Expert, France

10:30 COFFEE BREAK

11:00 Topic 1 Session 1 continued

BASIC DEFINITIONS AND TOOLS IN PHARMACOVIGILANCE
Sabine Brosch, European Medicines Agency, EU
Gaby Danan, Pharmacovigilance Expert, France

12:00 LUNCH

13:00 Topic 2 Session 2

PREPARATION OF DEVELOPMENT SAFETY UPDATE REPORTS (DSURs)
Vicki Edwards, AbbVie Ltd, UK

13:45 Topic 2 Session 3

PREPARATION OF PERIODIC SAFETY UPDATE REPORTS (PSURs)
Vicki Edwards, AbbVie Ltd, UK

14:30 Topic 2 Session 4

THE ROLE OF THE QUALIFIED PERSON RESPONSIBLE FOR PHARMACOVIGILANCE
Vicki Edwards, AbbVie Ltd, UK

15:15 COFFEE BREAK

15:30 Topic 2 Session 5

EXPEDITED REPORTING REQUIREMENTS IN THE POST-AUTHORIZATION PHASE AND CASE STUDIES
Sabine Brosch, European Medicines Agency, EU
Gaby Danan, Pharmacovigilance Expert, France

17:00 COFFEE BREAK

17:15 Topic 2 Session 5 continued

EXPEDITED REPORTING REQUIREMENTS IN THE POST-AUTHORIZATION PHASE AND CASE STUDIES
Sabine Brosch, European Medicines Agency, EU
Gaby Danan, Pharmacovigilance Expert, France

18:15 END OF DAY ONE

Tuesday | 14 October 2014

08:30 TOPIC 2

REGULATORY ASPECTS IN PHARMACOVIGILANCE AND PRACTICAL EXAMPLES
The roles and responsibilities of marketing authorisation holders and national Competent Authorities in the conduct of Pharmacovigilance are defined in EU legislation and further detailed in the Good Pharmacovigilance Practices (GVP). Topic 2 will provide a concise summary of the individual and periodic adverse reaction reporting requirements of marketing authorisation holders in the post-authorisation phase and illustrations based on practical case studies.

Furthermore, the roles and responsibilities of all stakeholders of interventional clinical trials, in line with the implementing texts published in relation to Directive 2001/20/EC, are summarised.

Aspects that need to be taken into account in establishing a Pharmacovigilance database as well as the key functionalities of the EU’s EudraVigilance system will be discussed.

08:30 Topic 2 Session 1

SUSAR REPORTING IN INTERVENTIONAL CLINICAL TRIALS AND CASE STUDIES
Sabine Brosch, European Medicines Agency, EU
Gaby Danan, Pharmacovigilance Expert, France

10:00 COFFEE BREAK

10:30 Topic 2 Session 1 continued

SUSAR REPORTING IN INTERVENTIONAL CLINICAL TRIALS AND CASE STUDIES
Sabine Brosch, European Medicines Agency, EU
Gaby Danan, Pharmacovigilance Expert, France

12:00 LUNCH

13:00 Topic 2 Session 2

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Vicki Edwards, AbbVie Ltd, UK

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15:15 COFFEE BREAK

15:30 Topic 2 Session 5

EXPEDITED REPORTING REQUIREMENTS IN THE POST-AUTHORIZATION PHASE AND CASE STUDIES
Sabine Brosch, European Medicines Agency, EU
Gaby Danan, Pharmacovigilance Expert, France

17:00 COFFEE BREAK

17:15 Topic 2 Session 5 continued

EXPEDITED REPORTING REQUIREMENTS IN THE POST-AUTHORIZATION PHASE AND CASE STUDIES
Sabine Brosch, European Medicines Agency, EU
Gaby Danan, Pharmacovigilance Expert, France

18:15 END OF DAY TWO

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TRAINING COURSE LOCATION
European Medicines Agency
Churchill Place 30
Canary Wharf
London
United Kingdom

Please see below website for conveniently located hotels:
http://www.londontown.com/hotels/
WEDNESDAY | 15 OCTOBER 2014

08:30  Topic 2 Session 5 continued

EXPEDITED REPORTING REQUIREMENTS IN THE POST-AUTHORISATION PHASE AND CASE STUDIES
Sabine Brosch, European Medicines Agency, EU
Gaby Danan, Pharmacovigilance Expert, France

10:30  COFFEE BREAK

11:00  Topic 2 Session 6

REPORTING REQUIREMENTS IN SPECIAL SITUATIONS IN THE POST-AUTHORISATION PHASE AND CASE STUDIES
William Gregory, Pfizer, USA

12:30  LUNCH

13:30  Topic 2 Session 7

PHARMACOVIGILANCE SYSTEM MASTER FILE (PSMF)
Sabine Brosch, European Medicines Agency, EU
Jerome Calmejane, Janssen-Cilag, France

14:30  COFFEE BREAK

14:45  Topic 2 Session 7 continued

PHARMACOVIGILANCE SYSTEM MASTER FILE (PSMF)
Sabine Brosch, European Medicines Agency, EU
Jerome Calmejane, Janssen-Cilag, France

15:45  COFFEE BREAK

16:00  Topic 2 Session 7 continued

AUDITS AND INSPECTIONS IN PHARMACOVIGILANCE
Jerome Calmejane, Janssen-Cilag, France

18:00  END OF DAY THREE

THURSDAY | 16 OCTOBER 2014

08:30  TOPIC 3

DIAGNOSIS AND MANAGEMENT OF ADVERSE DRUG REACTIONS
Pharmacovigilance is first based on the medical assessment of the adverse events passively or actively collected in organised schemes. It is then essential to be able to identify consistently the nature of events, their seriousness, their likelihood of occurrence and to assess causality with the suspect drug(s). This session will provide clues for the recognition of two serious events involving target organs of drug toxicity.

08:30  Topic 3 Session 1

MEDICAL EVALUATION OF ADVERSE DRUG REACTIONS
Gaby Danan, Pharmacovigilance Expert, France

09:30  Topic 3 Session 2

DRUG-INDUCED LIVER INJURY
Gaby Danan, Pharmacovigilance Expert, France

09:30  Topic 3 Session 2

DRUG-INDUCED LIVER INJURY
Gaby Danan, Pharmacovigilance Expert, France

10:30  COFFEE BREAK

11:00  Topic 3 Session 2 continued

DRUG-INDUCED LIVER INJURY
Gaby Danan, Pharmacovigilance Expert, France

11:30  Topic 3 Session 3

QT/QTc PROLONGATION AND THE RISK OF TORSADE DE POINTES
Gaby Danan, Pharmacovigilance Expert, France

12:30  LUNCH

13:30  TOPIC 4

SIGNAL DETECTION
New safety signals may emerge at any time following product launch and must be evaluated for relative risk, medical importance, and likelihood of occurrence. This session will provide an understanding of safety data classification, using MedDRA terminology and Standardised MedDRA Queries (SMQs) and approaches to signal detection using traditional and quantitative methods.

13:30  Topic 4 Session 1

MedDRA AND STANDARDISED MedDRA QUERIES (SMQs)
William Gregory, Pfizer, USA

14:15  Topic 4 Session 2

INTRODUCTION TO SIGNAL DETECTION
Georgy Genov, European Medicines Agency, EU

15:00  COFFEE BREAK

15:30  Topic 4 Session 3

SIGNAL MANAGEMENT IN THE EUROPEAN UNION
• Regulatory Network Perspective
  Georgy Genov, European Medicines Agency, EU
• Industry Perspective
  Jan Petracek, PharmInvent, Czech Republic

16:30  TOPIC 5

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 epidemiological methods.

This session aims to provide the background for understanding drug-related risks, to review epidemiological methods for detecting signals and assessing risks, and to present recent developments regarding risk communication.

16:30  Topic 5 Session 1

RISK COMMUNICATION IN EU - CHALLENGES AND POSSIBILITIES
Jan Petracek, PharmInvent, Czech Republic

18:00  END OF DAY FOUR

FRIDAY | 17 OCTOBER 2014

08:30  Topic 5 Session 2

RISK MANAGEMENT COMPONENTS: GENERAL PRINCIPLES
Thomas Goedecke, European Medicines Agency, EU

09:45  Topic 5 Session 3

RISK MANAGEMENT PLANS: AN INDUSTRY PERSPECTIVE
Jan Petracek, PharmInvent, Czech Republic

10:45  COFFEE BREAK

11:15  Topic 5 Session 4

POST-AUTHORISATION DEVELOPMENT PLAN (PASS/PAES)
Xavier Kurz, European Medicines Agency, EU

11:45  Topic 5 Session 5

EFFECTIVENESS OF RISK MINIMISATION MEASURES
Jan Petracek, PharmInvent, Czech Republic

12:45  END OF TRAINING COURSE
REGISTRATION FORM
EMA Excellence in Pharmacovigilance: Clinical trials and post-marketing training course
13-17 October 2014 | Churchill Place 30, Canary Wharf, London, UK

FEES

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Please complete in block capital letters or attach the attendee’s business card here.

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All cancellations must be made in writing and be received at the DIA Europe office five working days prior to the event start date. Cancellations are subject to an administrative fee:

• Industry (Member/Non-member) € 200.00
• Academy/Charitable/Government/Non-profit (Full-time) (Member/Non-member) € 100.00
• Tutorial cancellation € 50.00

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