eXtended EudraVigilance Medicinal Product Dictionary Training Course

Course #14159 9-10 October 2014 AEMPS, Madrid, Spain



Key Topics

- General Terms and Definitions
- Registration in EudraVigilance and Qualified Person Responsible for Pharmacovigilance (QPPV) registration (incl. sponsor registration)
- XEVPRM XSD Schema
- XEVPRM data elements and examples including hands-on exercises
- Operation Types
- Data Quality
- Data Ownership
- XEVMPD technical validation rules
- Use of Controlled Vocabularies

Course Goals

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

- Understand the concepts related to the electronic submission of information on medicines authorised in the ELL.
- Describe the format and the data elements of the XEVPRM for authorised medicinal products
- Discuss practical examples of different types of medicinal products
- Get hands-on experience in working with the XE-VMPD
- Describe the format and the data elements of the XEVPRM for IMPs

Details of the face-to-face training courses:

Duration: 2 days Location: AEMPS

Parque Empresarial Las Mercedes

Edificio 8, 3ª planta-A c/ Campezo, 1 E-28022-Madrid

The course is limited to 16 participants. Register early! Training on electronic submission of information on medicines New pharmacovigilance legislation (Art. 57, paragraph 2, 2nd sub-paragraph, Regulation (EC) No. 726/2004)

Introduction

The European Medicines Agency (EMA) is implementing the electronic submission of information on medicines in the context of the new pharmacovigilance legislation (Art. 57, paragraph 2, 2nd subparagraph,Regulation (EC) No. 726/2004). On 05 March 2012, EMA published an updated set of mandatory requirements for marketing authorisation holders to comply with Article 57(2). The number of data fields initially required in the format published on 2 July 2011 was considerably reduced, thus significantly reducing the administrative burden and helping marketing authorisation holders to meet their legal deadline of 2 July 2012.

With regard to investigational medicinal products (IMPs), EMA is also facilitating the implementation of the provisions set out in the detailed guidance on the collection, verification and presentation of adverse event/reaction reports arising from clinical trials on medicinal products for human use ("CT-3", chapter 7.9, paragraph 104).

Course Overview

The EMA has prepared this eXtended EudraVigilance Medicinal Product Dictionary (XEVMPD) course to facilitate the practical implementation of the requirements including technical aspects and all related procedures on electronic submission of information on medicines by marketing authorisation holders in the European Union (EU).

The training focuses on explaining the guidance, specifically the mandatory data elements necessary for the electronic submission of information on medicinal products applying the format of the eXtended EudraVigilance Product Report Message (XEVPRM) and the use of the XEVMPD data entry tool also known as EVWEB.

Participants who successfully pass the knowledge evaluation following the course will receive a notification from the European Medicines Agency that will allow them to register with EudraVigilance for the electronic submission of information on medicines in accordance with Article 57(2), second subparagraph of Regulation (EC) No. 726/2004.

The course also includes instructions for sponsors of clinical trials as how to provide information on the IMPs in the EudraVigilance Medicinal Product Dictionary ('EVMPD') before completing the clinical trials application form.

Course Audience

The XEVMPD training programme is targeting personnel of marketing authorisation holders, consultants and other organisations, who are responsible for the electronic submission and maintenance of information on medicinal products authorised in the EU.

It is also targeting sponsors of clinical trials responsible for providing information on IMPs in accordance with the CT-3 detailed guideline.

The content of this training course is subject to regular updates in order to comply with new regulations and requirements.

http://www.ema.europa.eu/ema/index.jsp?curl=pages/regulation/general/general_content_000492.jsp&murl=menus/regulations/regulations.isp&mid=WC0b01ac058033e8ad&isenabled=true







What this course offers

- Training in meeting the requirements of the provisions of Article 57(2), second sub-paragraph of Regulation (EC) 726/2004 and the electronic submission of
 information on authorised medicinal products
- Training in supporting the electronic submission of information on authorised medicinal products for Gateway users
- Training in developing messages compliant with the published XEVPRM XSD schemas
- Training in supporting the electronic submission of information on authorised medicinal products for Web trader and XEVMPD users
- Hands-on training using the XEVMPD to generate XEVPRMs
- Training in meeting the requirements of the provisions set out in the detailed guidance ("CT-3") and the electronic submission of information on IMPs

What this course does not cover

- Training in developing and validating information or communication technology tools to produce messages compliant with the published XEVPRM and SSI XSD schemas
- Training on all five ISO Identification of Medicinal Products (IDMP) standards and the Individual Case Safety Report (ICSR) standard as well as related ICH Implementation Guides
- Training on IDMP, ICSR and Common Product Model (CPM) HL7 V3 messages

Course Pre-requisites

Participants are expected to have basic background knowledge of:

- EU legislation and the revised guidance on the electronic submission of information on medicinal products for human use by marketing authorisation holders to the European Medicines Agency in accordance with Article 57 (2), second subparagraph of Regulation (EC) 726/2004
- Detailed guidance on the collection, verification and presentation of adverse event/reaction reports arising from clinical trials on medicinal products for human use ("CT-3", cahpter 7.9, pragraph 104).

Hotel and Travel Information

Attendees must make their own bookings. Recommended hotels nearby:

AXOR BARAJAS HOTEL Calle Campezo 4 Madrid 28022 Tel. +34 91 312 1960

E-mail: reservas.barajas@axorhoteles.com http://en.axorhoteles.com/suites-barajas/

HOTEL HILTON MADRID AIRPORT Av. de la Hispanidad 2 - 4 Madrid 28042, Tel. +34 91 153 4000

HOTEL NH BARAJAS Catamarán, 1 28042 Barajas, Madrid Tel. +34 91 742 0200 www.nh-hotels.com

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DIA Upcoming Training Courses in Safety and Pharmacovigilance

■ Benefit/Risk Management

May 2014 | Location tbc | More information available soon October 2014 | Location tbc | More information available soon

- Diagnosis and Management of Drug-Induced Liver Injury (DILI)
 September 2014 | Location tbc | More information available soon
- How to Prepare for Pharmacovigilance Audits and Inspections November 2014 | Location tbc | More information available soon
- ICH Endorsed Pharmacovigilance

Pre-Marketing Clinical Safety

Next recurrence of this course to be announced

June 2014 | Location tbc | More information available soon

Signal Management in Pharmacovigilance

May 2014 | Location tbc | More information available soon November 2014 | Location tbc | More information available soon

European Medicines Agency Information Days and Courses

- Excellence in Pharmacovigilance: Clinical trials and post-marketing 17-21 February 2014 | London, United Kingdom | ID 14500
- MedDRA Information Day

Next recurrence of this course to be announced

■ EudraVigilance Information Day

Next recurrence of this course to be announced

- EudraVigilance courses:
 - EudraVigilance Electronic reporting of ICSRs in the EEA
 - eXtended EudraVigilance Medicinal Product Dictionary
 - Introduction to Pharmacovigilance and Electronic Transmission of Individual Case Safety Reports (ICSR) for the Use of Eudravigilance at the European Medicines Agency

Courses throughout the year | European Medicines Agency, London, United Kingdom and selected European cities.

More information on EudraVigilance courses will be available soon, please visit www. diahome.org > Meetings & Training > About Meetings & Training > In-Person Instruction > EudraVigilance > EudraVigilance Courses.

Course Agenda

DAY ONE

8:45 Session 1

Course Introduction

Introduction to EudraVigilance

Registration to EudraVigilance

Session 2

Regulatory Background

General Terms and Definitions

eXtended EudraVigilance Medicinal Product Report Message (XEVMPRM) Data Set

Operation Types

Data Quality

Data Ownership

Session 3

Database Architecture

Roles of the eXtended EudraVigilance Medicinal Product Dictionary (XEVMPD) within EudraVigilance

Data Collection Process

COFFEE BREAK

Session 4

How to enter product data in the XEVMPD using the EVWEB tool

How to enter an organisation (MAH and Sponsor)

How to enter a substance (an approved and a development substance), translations and synonyms

LUNCH

Session 4 continued

Examples of different types of authorised medicinal products

- · Nationally authorised medicinal product
- · Medicinal product authorised through the mutual recognition procedure
- · Centrally authorised medicinal product

Investigational Medicinal Product (Development Medicinal Product) for sponsors of clinical trials

DAY TWO

8:45 Session 5

How to perform simple and advanced queries in the XEVMPD using the EudraVigilance Web-based application (EVWEB)

Session 6

How to maintain product data in the XEVMPD using EVWEB How to use the operation type "withdraw" for an authorised medicinal product

COFFEE BREAK

Example how to use the operation type "update" for substance (including the handling of translations and synonyms)

Example how to use the operation type "update" for an organisation

SANDWICH LUNCH

Knowledge Evaluation

Part 1: Multiple Choice Questions Part 2: Product Report Exam Case

17:00 END OF DAY TWO

The Agenda is subject to change as course content is updated regularly in order to comply with new regulations and requirements.

REGISTRATION FORM

eXtended EudraVigilance Medicinal Product Dictionary Training Course Course #14159 | 9-10 October 2014 | Madrid, Spain



The registration fee includes training course material, IT equipment, lunches and refreshments.

FAX YOUR COMPLETED REGISTRATION FORM TO: +41 61 225 51 52 or email to: diaeurope@diaeurope.org

FEES	The registration fee includes training course material, IT equipment, lunches and refreshments.
Standard Fee* € 1'180.00 □	TOTAL AMOUNT DUE:
Reduced Fee for Academia/Non-profit (Full-time)* € 585.00 □	
Reduced Fee for Government* € 585.00 □	Each course is limited to 16 participants. Courses may be cancelled if numbers of participants are not sufficient.
Please advise your Spanish VAT number:	Payment of registration fees must be received before commencement of the course.
*All fees are subject to Spanish VAT at 21 %.	
Special discount - for SME (status confirmed by EMA) available. Multiple course discount available if booked together with the three day EudraVigilance training course.	
ATTENDEE DETAILS	PAYMENT METHODS
PLEASE COMPLETE IN BLOCK CAPITAL LETTERS OR ATTACH THE ATTENDEE'S BUSINESS CARD HERE	Credit cards: Payments by VISA, Mastercard or AMEX can be made by completing the details below. Please note that other types of credit card cannot be accepted.
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DIA reserves the right to include your name and affiliation on the attendee list.	

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 € 200.00.
- Academia/Charitable/Government /Non-profit (Full-time) € 100.00.

If you do not cancel five working days prior to the event start date and do not attend, you will be responsible for the full registration fee. DIA Europe reserves the right to alter the venue and dates if necessary. If an event is cancelled DIA Europe is not responsible for airfare, hotel or other costs incurred by registered attendees. Registered attendees are responsible for cancelling their own hotel and travel reservations.

Transfer Policy

You may transfer your registration to a colleague prior to the start of the event. Please notify the DIA Europe office of any such substitutions as soon as possible.