

# ICH E2B (R3) Individual Case Safety Report (ICSR) Information Day

13 May 2014  
Course #14502  
European Medicines Agency (EMA), London, UK



## Programme Committee

**Paolo Alcini**  
Head Data Collection and Management, European Medicines Agency (EMA), EU

**Peter Richard Arlett**  
Head of Pharmacovigilance, European Medicines Agency (EMA), EU

**Sabine Brosch**  
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**Gaby Danan**  
Pharmacovigilance Expert, France

**Anja van Haren**  
EudraVigilance Coordinator, Medicines Evaluation Board (MEB), The Netherlands

## Details of the Information Day

Location: European Medicines Agency  
Canary Wharf  
7 Westferry Circus  
London E14 4HB, UK

Capacity: The event is limited to 120 participants

## Overview

In November 2012, step 4 of the ICH E2B (R3) package has been signed off based on the ISO ICSR standard including the awaited implementation guide (IG) accompanied by several technical appendices. This step opened the way for the worldwide implementation of the ISO ICSR standard replacing progressively the current E2B (R2) version. The first package (version 1.01) was made available on 12 April 2013 to the users in order to begin the testing phase and the implementation of data exchange between partners.

In the context of the EU implementation, a regional IG is being prepared addressing EU specific requirements in relation to the application of the ISO ICSR standard and the E2B (R3) package.

This Information Day will address and explain the key changes expected in relation to the application of the new ISO ICSR standard and how those will impact the EU adverse reaction reporting and electronic transmission activities.

## Key Topics

- Key differences between the ISO ICSR International Standard and the current ICH E2B(R2) guideline
- The ICH safety message flow in the EU
- Processing of safety and acknowledgement messages in case of technical or system failures
- EU specific business rules and technical ICSR validation
- Case classification
- ICSR specific concepts and their application in the EU (e.g. amendment report, causality assessment)
- Coding of medicinal product information
- Use of MedDRA in the context of the new ICSR reporting
- Handling of attachments
- EMA testing procedures with stakeholders

## Learning Objectives

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

- Recognise the new requirements as regards the ICH E2B (R3) and EU region specific implementation
- Prepare for the implementation of the new ICSR standard and the adaptation of internal pharmacovigilance systems by all stakeholders involved (medicines regulatory authorities in the EU, IT vendors and pharmaceutical companies)
- Understand the use of the new ICSR format in line with EU pharmacovigilance legislation

## Who Will Attend

- Representatives of IT departments of medicines regulatory authorities, pharmaceutical companies and service providers
- EU Qualified Persons Responsible for Pharmacovigilance (EU QPPVs)
- Pharmacovigilance staff of pharmaceutical companies and medicines regulatory authorities
- Pharmacovigilance software vendors
- Sponsors of Clinical Trials

**8:45 Welcome and Opening Remarks**

Peter Richard Arlett, EMA, EU

Session chairs:

Anja van Haren, MEB, NL and Sabine Brosch, EMA, EU

**9:00 Session 1****KEY DIFFERENCES BETWEEN THE NEW E2B (R3) ICSR AND THE ICH ICSR E2B (R2)**

This session will provide a summary of the differences between the new ICH E2B (R3) and the current E2B (R2) ICSR format in the context of the electronic reporting of adverse reactions in the EU. The expected benefits and the impact on the pharmacovigilance business processes will be highlighted.

**Speakers:**

Anja van Haren, MEB, NL

Gaby L. Danan, Pharmacovigilance Expert, France

Discussant: Diane Farkas, Sanofi-Aventis, France

**10:00 Session 2****ELECTRONIC ICSR REPORTING PROCESS**

This session will describe the procedures concerning the Electronic Data Interchange (EDI) of ICSRs and the roles of all involved stakeholders taking into account the simplification of adverse reaction reporting as foreseen in Article 107(3) of Directive 2001/83/EC and Article 28(1) of Regulation (EC) 726/2004.

The ICSR safety message flow in the EU

Nick Halsey, EMA, EU

**10:45 Coffee Break****11:15 Session 3****EU SPECIFIC BUSINESS RULES AND TECHNICAL ICSR VALIDATION**

Key changes to the business rules as currently applied in EudraVigilance will be presented. These changes are based on the new ISO ICSR standard, the ICH E2B (R3) Implementation Guide and taking into account EU specific requirements and processes.

EU specific business rules and case classification

Nick Halsey, EMA, EU and Edurne Lazaro, AEMPS, ES

**12:30 Sandwich lunch****13:30 Session 4****ICSR SPECIFIC CONCEPTS AND THEIR APPLICATION IN THE EU**

This session will address the handling of amendment reports, attachments and principles of causality of assessment. Principles of handling medicinal product information will be also elaborated.

Concepts of the new ICSR applied in the EU

Anja van Haren, MEB, NL

Discussants: Sabine Brosch, EMA, EU and Victoria Newbould, EMA, EU

Handling of medicinal product information in ICSRs

Tom Paternoster-Howe, EMA, EU and Ilaria Del Seppia, EMA, EU

Discussant: Ana Silvia Cochino, EMA, EU

**15:00 Coffee Break****15:30 Session 5****EMA TESTING PROCEDURES AND INDUSTRY PERSPECTIVES**

The Preparation for the new ICSR implementation from a pharmaceutical industry perspective

Diane Farkas, Sanofi-Aventis, France

An outline of potential testing procedures for the new E2B (R3) ICSR format

Tom Paternoster-Howe, EMA, EU

**16:45 END OF INFORMATION DAY****ABOUT DIA**

The DIA is a global association of approximately 18,000 members who are involved in the discovery, development, regulation, surveillance or marketing of pharmaceuticals or related products. The DIA is committed to the broad dissemination of information on the development of new medicines or generics, biosimilars, medical devices and combination products with continuously improved professional practice as the goal.

The DIA is an independent non-profit organisation. The voluntary efforts of DIA members and speakers allow the DIA to organise conferences, workshops and training courses and provide publications and educational material.

DIA's headquarters are in Horsham, PA, USA, with the European office in Basel, Switzerland, and other regional offices in Tokyo, Japan, Mumbai, India, and Beijing, China.

For more information, visit [www.diahome.org](http://www.diahome.org) or call DIA Europe on +41 61 225 51 51.

Unless otherwise disclosed, DIA Europe acknowledges that the statements made by speakers are their own opinion and not necessarily that of the organisation they represent, or that of the DIA Europe.

Speakers and agenda are subject to change without notice. Recording of any DIA Europe tutorial/workshop information in any type of media, is prohibited without prior written consent from DIA Europe.

## HOTEL INFORMATION

### Recommended Hotel:

Hilton London Docklands Riverside

265 Rotherhithe Street, London, SE16 5HW, UK

Telephone: +44 20 7231 1001

Email: [reservations.docklands@hilton.com](mailto:reservations.docklands@hilton.com)

DIA has blocked a limited number of rooms at the rate of GBP 139.00 single and GBP 149.00 double room/night including breakfast and VAT. To make your booking please visit the DIA event website.

The hotel is situated opposite of Canary Wharf, conveniently connected by a shuttle boat. The landing stage is in walking distance to the European Medicines Agency (2 min). The ferry ticket is included in the room rate. Please make sure you receive it when checking in.

For further information, please go to [http://www1.hilton.com/en\\_US/hi/hotel/LONNDHI-Hilton-London-Docklands-hotel/index.do](http://www1.hilton.com/en_US/hi/hotel/LONNDHI-Hilton-London-Docklands-hotel/index.do)

## DIA 2014 Training Courses in Safety and Pharmacovigilance

- **Benefit/Risk Management**  
19-20 May 2014 | Prague, Czech Republic | ID 14533
- **Signal Management in Pharmacovigilance**  
21-22 May 2014 | Prague, Czech Republic | ID 14534
- **Pre-Marketing Clinical Safety**  
16-17 June 2014 | Location to be confirmed | ID 14539
- **Post-Authorisation Safety Studies (PASS) new offering!**  
18-19 June 2014 | Location to be confirmed | ID 14535
- **Medical Approach in Diagnosis and Management of ADRs**  
22-23 September 2014 | Paris, France | ID 14544
- **Diagnosis and Management of Drug-Induced Liver Injury (DILI)**  
23-24 September 2014 | Paris, France | ID 14544
- **ICH Endorsed Pharmacovigilance**  
21 October 2014 | Dakar, Senegal | ID 14559
- **Benefit/Risk Management**  
October 2014 | Location to be confirmed | ID 14547
- **How to Prepare for Pharmacovigilance Audits and Inspections**  
November 2014 | Location to be confirmed | ID 14550
- **Diagnosis and Management of Drug-Induced Liver Injury (DILI)**  
November 2014 | Algiers, Algeria | ID 14560
- **Signal Management in Pharmacovigilance**  
November 2014 | Location to be confirmed | ID 14549

## European Medicines Agency Information Days and Courses

- **ICSR Information Day**  
13 May 2014 | London, United Kingdom | ID 14502
  - **EnCePP Information Day**  
Dates to be confirmed | London, United Kingdom | ID 14503
  - **Excellence in Pharmacovigilance: Clinical trials and post-marketing**  
13-17 October 2014 | London, United Kingdom | ID 14548
  - **MedDRA Information Day**  
November 2014 | London, United Kingdom | ID 14549
- EudraVigilance courses:
- EudraVigilance – Electronic reporting of ICSRs in the EEA
  - eXtended EudraVigilance Medicinal Product Dictionary
  - Introduction to Pharmacovigilance and Rules for Expedited Reporting of Individual Case Safety Reports (ICSRs) in Europe

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

For information on EudraVigilance courses, please visit [www.diahome.org](http://www.diahome.org) > Meetings & Training > About Meetings & Training > In-Person Instruction > EudraVigilance > EudraVigilance Courses

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# REGISTRATION FORM

6th ICH E2B (R3) Individual Case Safety Report (ICSR) Information Day  
Course #14502 | European Medicines Agency (EMA), London, UK



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

## FEES

Standard Fee	€	365.00	<input type="checkbox"/>
Reduced Fee for Academia/Government/ Non-profit (Full-Time)	€	150.00	<input type="checkbox"/>

The registration fee includes training course material, sandwich lunch and refreshments.

TOTAL AMOUNT DUE: \_\_\_\_\_  
Payment of registration fees must be received before commencement of the course.

## ATTENDEE DETAILS

PLEASE COMPLETE IN BLOCK CAPITAL LETTERS OR ATTACH THE ATTENDEE'S BUSINESS CARD HERE

Prof  Dr  Ms  Mr

Last Name

First Name

Company

Job Title

Address

Postal Code  City

Country

Telephone

Fax

Email\*

\*(Required for confirmation)

DIA reserves the right to include your name and affiliation on the attendee list.

## PAYMENT METHODS

**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.

Please charge my  VISA  MC  AMEX

Card N°

Exp. Date  /

Cardholder's Name

**Bank transfers:** When DIA completes your registration, an email will be sent to the address on the registration form with instructions on how to complete the bank transfer. Payments in EURO should be addressed to "Account Holder: DIA." Please include your name, company, Event ID# 14502 as well as the invoice number to ensure correct allocation of your payment.

Payments must be net of all charges and bank charges must be borne by the payer. **If you have not received your confirmation within five working days, please contact DIA Europe.**

If you require a hardcopy of our terms and conditions, please contact our Customer Service Team.

By signing below, I confirm that I agree with DIA Europe's Terms and Conditions of booking.

Date

Signature

## Cancellation Policy

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:

- Full Meeting Cancellation: Industry (Member/Non-member) = € 200.00.
- Academia/Charitable/Government /Non-profit (Full-Time) (Member/Non-member) = € 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.