

DIA/MEB Excellence in Pharmacovigilance: Good Pharmacovigilance Practices (GVP) Modules I to XVI

14-17 November 2022
Amsterdam, The Netherlands*

OVERVIEW

Organized and delivered in collaboration with the Dutch Medicines Evaluation Board (MEB), this face-to-face training course covers the major pharmacovigilance processes as outlined in GVP I to XVI. It is designed to provide a strong foundation in all key aspects of European Post-Marketing Safety regulatory requirements.

Furthermore, it includes highlights and updates on the pharmacovigilance legislation as well as the latest news on the ICH activities in pharmacovigilance.

This training course will take place in Amsterdam, allowing live interaction and real-time Q&A with the large panel of renowned experts onsite. We have also decided to make this course available for virtual participation, with some limitation regarding interaction with fellow participants and faculty members. You can register for your preferred format immediately and without concern: DIA will monitor the situation and keep all registrants informed on a regular basis.

LEARNING OBJECTIVES

Additionally to the key topics as outlined below, the learning objectives also include the ability to:

- Understand basic definitions and methods in Pharmacovigilance
- Describe the expedited and periodic ICSRs reporting requirements in post-marketing
- Understand the process of audits and inspections in pharmacovigilance
- Understand the principles of signal management
- Describe the components of the risk management

KEY TOPICS

- Post-Marketing: GVP Modules VI and VII
- Regulatory Aspects in Pharmacovigilance: GVP Modules I, II, III and IV
- Risk Management: GVP Modules V, VIII, XV and XVI
- Signal Management: GVP Modules IX and X

TARGET AUDIENCE

Professionals with experience in safety related activities of the drug development process and/or those with a need of a holistic overview about all PV related regulatory requirements will benefit most, such as:

- Pharmacovigilance Officers, Managers, Specialists, Experts, or Coordinators
- Regulatory Compliance, Quality or Safety departments Heads, Directors or Managers

Level: /Intermediate

**Please note that in case the epidemiological situation does not allow a face-to-face course, this offering will take place in a virtual setting.*



COURSE DIRECTORS

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DIA

C B G
M E B

PharmaTrain
MASTERING MEDICINES DEVELOPMENT

14 NOV | DAY 1 | POST-MARKETING

Day 1 corresponds to GVP Modules VI and VII and will cover individual and periodic adverse reaction reporting requirements of marketing authorisation holders in the post-authorisation phase with illustrations based on case studies as practical examples.

08:00 Registration**08:30 Welcome and Introduction****08:45 Keynote: PRAC: Present and Future of Pharmacovigilance**
Liana Gross-Martirosyan**09:30 COFFEE BREAK****10:00 Expedited Reporting Requirements in the Post-Authorisation Phase and Case Studies**
Wendy Huisman**12:30 LUNCH BREAK****13:30 Preparation of Aggregate Reports (PSUR and DSUR)**
Mark Bailey**15:00 COFFEE BREAK****15:30 Reporting Requirements in Special Situations in the Post-Authorisation Phase**
Wendy Huisman**16:30 Discussion and Q&A****17:00 WELCOME RECEPTION****18:00 END OF DAY 1****| Event Safety Code of Conduct**

In 2022, DIA events are returning to a face-to-face format. Because we are committed to delivering a safe and secure conference, please be advised that all in-person participants must comply with our [Event Safety Code of Conduct](#), which requires either vaccination, certificate of recovery, or proof of inability to be vaccinated alongside a valid PCR test. Please consult the full [Event Safety Code of Conduct](#) for more information.

Unless otherwise disclosed, DIA 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. Speakers and agenda are subject to change without notice. Recording during DIA sessions is strictly prohibited without prior written consent from DIA.

15 NOV | DAY 2 | REGULATORY ASPECTS IN PV

Day 2 covers GVP Modules I to IV and will provide the safety reporting requirements with case studies, as well as safety data classification, using MedDRA terminology and safety data retrieval using Standardised MedDRA Queries (SMQs).

It covers establishment of a Quality Management System in Pharmacovigilance and includes aspects of the applicable GVP modules, as well as preparation and conduct of audits and inspections.

08:30 SUSAR Reporting in Clinical Trials and Case Studies
Wendy Huisman**10:45 COFFEE BREAK****11:15 MedDRA and Standardised MedDRA Queries**
Evelyn Mulder-Olthof**12:15 Pharmacovigilance System Master File**
Wendy Huisman**12:45 LUNCH BREAK****13:45 The Role of the Qualified Person Responsible for PV**
Wendy Huisman**14:15 Audits and Inspections in Pharmacovigilance – Regulatory Perspective**
Marian Verbruggen**15:15 COFFEE BREAK****15:45 Quality Management System**
Stephanie Martin**17:45 Discussion and Q&A****18:15 END OF DAY 2****| Course Venue****Holiday Inn Amsterdam - Arena Towers**

Hoogoorddreef 66A

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Email: info.amsaa@hiex.nlWeb: <https://www.ihg.com/holidayinn/hotels/us/en/amsterdam/amsaa/hoteldetail>**How to get there**

On of the largest train stations of Amsterdam; Amsterdam Bijlmer Arena is located on 5 minute walk of the hotel. When leaving the train station walk along the Heineken Music Hall in south-east direction and you will almost already see the hotel in front of you. Estimated distance to the hotel: 0.25 MI/ 0.4 KM

Bedroom booking

DIA has booked a limited number of hotel rooms at the rate of EUR 149.00 per standard single occupancy room per night including breakfast and VAT, but excluding local city tax at 7% per room per night and EUR 3.00 per person per night.

If you would like to make a booking, please do so [here](#).

The room rate is available until 13 October 2022 or until the room block is sold-out, whichever comes first.

16 NOV | DAY 3 | RISK MANAGEMENT

Day 3 corresponds to GVP Modules V, VIII, XV and XVI. 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. Specific examples of data collection and analysis will be presented in this session. Besides, pharmaco-epidemiological studies, which are the fundamentals of “additional” Pharmacovigilance activities, are discussed. This session also presents recent developments regarding risk communication.

08:30 Risk Management Plans – Regulatory Perspective: GVP Module V Revision 2
Paul ten Berg

09:30 Risk Management Plans – Industry Perspective
Jan Petracek

10:30 COFFEE BREAK

11:00 Harmonisation of RMP (HaRP) in Europe
Paul ten Berg

11:30 LUNCH BREAK

12:30 Epidemiological Methods and Pharmacovigilance
Fakhredin Sayed Tabatabaei

14:00 Effectiveness of Risk Minimisation Measures – Regulatory Perspective
Anita Volkers

15:00 COFFEE BREAK

15:30 Effectiveness of Risk Minimisation Measures – Industry Perspective
Jan Petracek

16:30 Risk Communication in EU – Challenges and Possibilities
Jan Petracek

17:30 Discussion and Q&A

18:00 END OF DAY 3

17 NOV | DAY 4 | SIGNAL MANAGEMENT

Day 4 covers GVP Modules IX and X. 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.

08:30 Introduction to Signal Detection in the European Union – Regulatory Perspective
Negar Babae and Evelyn Mulder-Olthof

10:30 COFFEE BREAK

11:00 Signal Management in the European Union – Industry Perspective
Jan Petracek

12:00 LUNCH BREAK

13:00 Signal Management – Workshop
Negar Babae and Evelyn Mulder-Olthof

15:00 Discussion and Q&A

15:30 Closing Remarks

15:45 END OF THE TRAINING COURSE

Group Discount

Register 3 individuals from the same company to the same course and receive complimentary registration for a 4th!

- All 4 individuals must register and prepay at the same time – no exceptions
- DIA will apply the value of the lowest applicable fee to this complimentary registration; it does NOT include fees for optional events or DIA membership
- You may substitute group participants of the same membership status at any time; however, administrative fees may be incurred.
- Group registration is not available online and does not apply to the already discounted fees for industry, government or charitable nonprofit/academia

To take advantage of this offer, please print the registration form for EACH of the four registrants from your company. Include the names of all four group registrants on each of the forms and return them together via email to basel@diaglobal.org.

Continuing Education

The Swiss Association of Pharmaceutical Professionals (SwAPP) and the Swiss Society for Pharmaceutical Medicine (SGPM) have accredited this training course with 27.5 credits.



REGISTRATION FORM

Excellence in Pharmacovigilance # 22548

14-17 November 2022 | Amsterdam, The Netherlands*

REGISTRATION FEES

Registration fee includes full admission to the training course and electronic access to training course materials. **Please note that the full amount must be received by DIA by commencement of the course to get the electronic access to the material.** Please check:

FEES*	MEMBER EARLY-BIRD valid until 19 Sep	MEMBER valid from 20 Sep	NON-MEMBER
INDUSTRY	€ 2'340.00 <input type="checkbox"/>	€ 2'600.00 <input type="checkbox"/>	€ 2'785.00 <input type="checkbox"/>
ACADEMIA/CHARITABLE/GOVERNMENT/NON-PROFIT (FULL-TIME)	NA	€ 1'300.00 <input type="checkbox"/>	€ 1'485.00 <input type="checkbox"/>

A special discount for SMEs on the standard fee is available for a limited number of places. To prove your status as an SME, a confirmation of the European Medicines Agency is necessary. Please contact DIA for more information

*All fees are subject to the VAT if applicable.

Please enter your Company's VAT number: _____

If DIA cannot verify your membership upon receipt of registration form, you will be charged the non-member fee.

***Please note that in case the epidemiological situation does not allow a face-to-face course, this offering will take place in a virtual setting.**

DIA MEMBERSHIP

All nonmember fees include a one year DIA membership, at no additional cost. Explore membership benefits at [DIAglobal.org/Membership](https://www.diaglobal.org/Membership).

DIA membership will renew automatically at the end of the complimentary membership term, at the then current membership rates. You may cancel automatic membership renewal at any time by accessing your account online at [DIAglobal.org](https://www.diaglobal.org). If you would like to decline complimentary membership, please indicate your preference below.

I would like to decline a one year complimentary DIA membership.

The DIA Contact Centre Team will be pleased to assist you with your registration from Monday to Friday between 09:00 and 17:00 CET/CEST.

Tel. :+41 61 225 51 51 Fax: +41 61 225 51 52 Email: Basel@diaglobal.org

Mail: DIA, Kuchengasse 16, 4051 Basel, Switzerland Web: www.DIAglobal.org



TERMS AND CONDITIONS

Cancellation Policy

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

- Industry (Member/Non-member) € 200.00
- Academia/Charitable/Government/Non-profit (Full-time) (Member/Non-member) € 100.00

If you do not cancel four weeks prior to the event start date and do not attend, you will be responsible for the full registration fee.

DIA reserves the right to alter the venue and dates if necessary. If an event is cancelled or postponed, DIA 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 but membership is not transferable. Substitute attendees will be responsible for the non-member fee, if applicable. Please notify the DIA office of any such substitutions as soon as possible.

Event Stream and Recording

If you attend a DIA event, we make video and audio recordings of events (both face-to-face and online) that may include your participation in the event, including your image, questions and comments. To view our full photography and video recording policy, click <https://www.diaglobal.org/general/photography-policy>.

Privacy Policy

DIA respects the privacy of all of its members and customers. To view our privacy policy, click <https://www.diaglobal.org/about-us/privacy-policy>. You agree that your personal data will be transferred to DIA in the US.

ATTENDEE DETAILS

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

Prof Dr Ms Mr

Last Name

First Name

Job Title

Company

Address

Postal Code

City

Country

Telephone Number

Attendee email required for course material access

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, Course ID #22548 as well as the invoice number to ensure correct allocation of your payment.

Please note: if you register 7 days or less before the start of the course, it is not possible to settle the registration fee by bank transfer, but only by credit card. Thank you for your understanding and cooperation.

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.**

By signing below, I confirm that I agree with DIA's Terms and Conditions of booking. These are available from the office or on <http://www.diaglobal.org/EUTerms>

Date

Signature