



Overview

Using pharmacovigilance audit techniques allows a company to identify any existing gaps or risks in their systems and procedures. This allows them to define and establish priorities, ensuring brand protection and company compliance.

Participants will learn how to prepare for an audit and inspection to achieve best practices from the moment of facing the auditing/inspection visit notification to the moment of receiving the report and its conclusions.

Learning Objectives

At the conclusion of this virtual live training course, participants will be able to:

- Plan pharmacovigilance audits based on risk assessment
- Identify and address the different areas of a pharmacovigilance system through audits
- Conduct a pharmacovigilance audit
- Evaluate audit documentation
- Manage communication with difficult characters, situations with missing documentation and master extreme situations
- Handle disagreements on audit findings
- Identify and follow-up on corrective and preventive actions (CAPAs)
- Host and manage a pharmacovigilance inspection

Who Will Attend

Those professionals most likely to benefit from this course will have experience in:

- Pharmacovigilance
- Drug Safety
- Regulatory Affairs
- Quality Assurance
- Risk Management
- Medical Affairs

or holding similar positions within the industry.

A sound knowledge of Pharmacovigilance is a must. Practical experience in audits and inspections is desirable.

Faculty

Calin Lungu

CEO
Drug Development Consulting Services
Luxembourg

Diane Hallé

Director, QPPV Compliance and Quality
Interface
Moderna
France

Key Topics

- PV audits - QMS requirements from GVP
- PV audit planning
- Operating individual PV audits
- Affiliates and third parties
- Reconciliation process
- Computerised systems
- PV inspection readiness
- Management of PV inspection
- Management of post-PV inspection activities

Schedule-At-A-Glance

DAY 1

13:00 WELCOME AND INTRODUCTION OF FACULTY AND PARTICIPANTS

13:30 SESSION 1

PV AUDITS – QMS REQUIREMENTS FROM GVP

- EU QPPV
- PSMF
- KPI
- Contractual agreements
- Business continuity

15:00 BREAK

15:30 SESSION 1 CONTINUED

PV AUDITS – QMS REQUIREMENTS FROM GVP

17:00 Q&A

17:30 END OF DAY 1

DAY 2

13:00 SESSION 2

PV AUDIT PLANNING

- Identify the different areas of the PV system to be audited
- Building a strategic and tactical PV audit planning
 - Case study: Identification of “PV audit universe”
 - Case study: Building strategic/tactical audit planning

15:00 BREAK

15:30 SESSION 3

OPERATING INDIVIDUAL PV AUDITS

- General process (plan, prepare, conduct, report, follow-up and CAPA)

17:00 INTRODUCTION TO HOMEWORK FOR DAY 3

CASE STUDY ON AUDITS OF AFFILIATES AND THIRD PARTIES

17:30 END OF DAY 2

DAY 3

13:00 SESSION 4

KEY PV AUDIT AREA 1: AFFILIATES AND THIRD PARTIES

- Audit planning (risk assessment, resources, audit team)
- Preparation (documentation requested in advance)
- Documentation audit e.g., PSURs
 - Case study review: Audit of affiliates and third parties

15:00 BREAK

15:30 SESSION 5

KEY PV AUDIT AREA 2: RECONCILIATION PROCESS

- ICSR: Internal reconciliation and reconciliation with interfaces (medical information, complaints department)
- ICSR: Reconciliations with external entities (distributors, license partners, market research contractors, PSP services)
- Databases reconciliation: Pharmacovigilance or clinical databases

17:30 END OF DAY 3

DAY 4

13:00 RECAP AND Q&A

13:30 SESSION 6

KEY PV AUDIT AREA 3: COMPUTERISED SYSTEMS

- Principles and contents of validation dossier
- Validation team
- Risk analysis
- Design qualification
- IQ, OQ, PQ, PQ I & PQ II
- Validation report
- Maintaining the validated status of the database

15:00 BREAK

15:30 SESSION 7

PV INSPECTION READINESS

- Checking resources (staff preparation, room and logistics)
- Running mock interviews with key staff
- Review of procedures
- Tour of facilities
- Remote audits and inspections

17:00 INTRODUCTION TO HOMEWORK FOR DAY 5

EACH PARTICIPANT TO CHOOSE ONE CASE STUDY

- Case study on computerised systems or audits as preparation for a PV inspection
- Case study on inspection findings

17:30 END OF DAY 4

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.

DAY 5

13:00 REVIEW OF HOMEWORK CASE STUDIES

13:45 SESSION 8

MANAGEMENT OF PV INSPECTION

- Logistics (staff preparation, room, recording document requests etc.)
- Do's and Don'ts during the inspection
- Disagreement with findings
- Closing meeting

14:15 BREAK

14:45 SESSION 9

MANAGEMENT OF POST-PV INSPECTION ACTIVITIES

- Receiving inspection report
- Handling additional documents' requests post-inspection
- Answering to findings and CAPA
- Agreeing on timelines
- How to prepare for a re-inspection

17:30 END OF THE TRAINING COURSE

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The dedicated efforts of DIA staff, members and speakers enable DIA to provide a comprehensive catalogue of conferences, workshops, training courses, scientific publications and educational materials. DIA is a global community representing thousands of stakeholders working together to bring safe and effective products to patients.

DIA is an independent, non-profit organisation has its Global Center in Washington, DC, USA with the European office in Basel, Switzerland, and additional regional offices in Horsham, Pennsylvania, USA; Tokyo, Japan; Mumbai, India; and Beijing, China.



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To test your system compatibility, please click on the link: <https://diaglobal.zoom.us/test>

For further information on system requirements, please visit the website: <https://www.diaglobal.org/General/System-Requirements>



Continuing Education

All DIA training courses have been awarded a PharmaTrain Centre Recognition.

PharmaTrain Federation is a not for profit organisation that started its activities as an IMI (Innovative Medicines Initiative) European Project. Its mission is to drive implementation of globally recognized high-level standards for postgraduate education and training in Medicines Development. To that aim, the Federation is assessing Continuous Professional Development (CPD) Courses and Course Providers around the world that deserve recognition.



REGISTRATION FORM

Advanced PV Audits and Inspections | Virtual Live Training Course | # 27551
15-19 November 2027 | 13:00-17:30 CET

DIA LEARNING

REGISTRATION FEES

Registration fee includes full admission to virtual course, 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 18 Oct 2027	MEMBER valid from 19 Oct 2027	NON-MEMBER
INDUSTRY/ REPRESENTATIVE	€ 1780.00 <input type="checkbox"/>	€ 1980.00 <input type="checkbox"/>	€ 2240.00 <input type="checkbox"/>
ACADEMIA/CHARITABLE/GOVERNMENT/NON-PROFIT (FULL-TIME)	NA	€ 990.00 <input type="checkbox"/>	€ 1250.00 <input type="checkbox"/>

A special discount is available for organisations which are listed in the [EMA SME register](#).
Number of discounted seats are limited.

All registration fees are subject to 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.

DIA MEMBERSHIP

All nonmember fees include a one year DIA membership, at no additional cost. Explore membership benefits at [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](#). 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 CE(S)T. **Tel.** :+41 61 225 51 51

Email: Basel@DIAglobal.org **Mail:** DIA, KÜchengasse 16, 4051 Basel, Switzerland

Web: www.DIAglobal.org

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

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.

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Date	Signature
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