D LEARNING

Pharmacovigilance System Master File

26-27 November 2024 | 13:00-17:00 CET



Overview

This virtual live training course covers essential concepts and guidance about the Pharmacovigilance System Master File (PSMF).

This key document describes the company's pharmacovigilance system, supporting, and documenting its compliance with the requirements laid down in the EU legislation and is the first document requested by a Competent Authority in preparation of a pharmacovigilance inspection.

The entire course is in line with the guidelines on EU Good Pharmacovigilance Practices GVP Module II - Pharmacovigilance System Master File (rev. 2), Commission Implementing Regulation (EU) No. 520/2012, and relevant EMA guidelines.

Participants benefit from hands-on expertise on best practices shared by trainer with extensive experience regarding PSMF including the EU-QPPV perspective. Ample time is set aside for Q&A and interactive discussions.

Learning Objectives

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

- Identify the structure, sections, and annexes of the PSMF
- Recognize the importance of the PSMF in the Pharmacovigilance system of a pharmaceutical company
- Evaluate the interaction between Regulatory Affairs, Pharmacovigilance, and other departments with regards to the maintenance of the PSMF
- Apply the essential concepts and principles of the GVP Module II -Pharmacovigilance System Master File (rev. 2)
- Prepare and manage this document in their own organisation
- Discuss the regulatory expectations for this important document, common inspection findings and gaps
- List quality performance indicators for monitoring timely submissions of ICSRs, PSURs and safety variations

Key Topics

- GVP Module II Pharmacovigilance System Master File (rev. 2) guidance
- Creation, maintenance, and management of the PSMF
- Practical exercise on drafting a PSMF
- The PSMF as a quality document
- Regulatory expectations for the PSMF, including UK PSMF
- Practical exercise on PSMF after an inspection

Who Will Attend

This virtual live training course is designed for professionals working in:

- Pharmacovigilance (including EU QPPVs)
- Drug Safety and Risk Management
- Pharmacovigilance Consultancies and Service Providers
- Quality and Compliance

Faculty

Jose Alberto Ayala Ortiz

CEO PVpharm Spain

Marcela Fialova

COOiVigee Services Czech Republic

MHRA speaker invited

MHRA United Kingdom



DAY 1

13:00 WELCOME AND INTRODUCTION

13:30 SESSION 1

GVP MODULE II - PHARMACOVIGILANCE SYSTEM MASTER FILE (REV. 2) GUIDANCE

Jose Ortiz

- Objectives, location and registration
- Responsibilities
- · Information to be contained, sections
- Annex

15:00 **BREAK**

15:15 SESSION 2

CREATION, MAINTENANCE, AND MANAGEMENT OF THE **PSMF**

Jose Ortiz

- Processes and workflows
- Interaction with other departments
- · Change control, log book, versions and archiving

16:30 SESSION 3

PRACTICAL EXERCISE ON DRAFTING A PSMF Jose Ortiz

17:00 END OF DAY 1

DAY 2

13:00 SESSION 4

THE PSMF AS A QUALITY DOCUMENT

Jose Ortiz

- The PSMF in the QMS
- · Audits, inspections

14:30 BREAK

14:45 SESSION 5

REGULATORY EXPECTATIONS FOR THE PSMF

Jose Ortiz and MHRA speaker invited

- Regulatory expectations
- Globalization
- UK PSMF

16:30 SESSION 6

PRACTICAL EXERCISE ON PSMF AFTER AN INSPECTION Jose Ortiz

17:00 END OF THE TRAINING COURSE

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Group Discounts

Register 3 individuals from the same company for the same course and receive complimentary registration

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.

*Terms and Conditions apply. Please contact DIA EMEA office for more information.



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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



Technical Requirements

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

The Swiss Association of Pharmaceutical Professionals (SwAPP) and the Swiss Society for Pharmaceutical Medicine (SGPM) have accredited this training course with 6 credits.



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.

REGISTRATION FORM

PSMF Virtual Live Training Course # 24553 26-27 November 2024 | 13:00-17:00 CET



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 1 Oct 2024	MEMBER valid from 2 Oct 2024	NON- MEMBER
INDUSTRY/ REPRESENTATIVE	€ 900.00 □	€ 1′000.00 🗖	€ 1′260.00 🗖
ACADEMIA/CHARITABLE/GOVERNMENT/NON-PROFIT (FULL-TIME)	NA	€ 500.00 □	€ 760.00 🗖
A special discount is available for organisations which are listed in the EMA SMF register:			

All registration fees are subject to VAT if applicable.

Please enter your company's VAT number:

https://fmapps.ema.europa.eu/SME/. Number of discounted seats are limited.

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

Payment is due 30 days after registration and must be paid in full by commencement of the course.

DIA MEMBERSHIP

All nonmember fees include a one year DIA membership, at no additional cost. Explore membership benefits at DIAqlobal.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: <u>Basel@DIAglobal.org</u> Mail: DIA, Küchengasse 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

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