

# Signal Management in Pharmacovigilance

14-15 November 2018
Mercure Lyon Charpennes Hotel, Lyon, France



This course will teach basic concepts of signal detection and signal management and how to apply them within the participants' functions, including the data mining techniques for large volume ADR data analysis, relevant EMA guidelines as well as a future outlook.

The entire course is updated in line with the latest guidelines on EU Good Pharmacovigilance Practices (GVP): Module IX – Signal management, Commission Implementing Regulation (EU) No. 520/2012, and CIOMS VIII. Time has been set aside for exercises, questions and discussions.

Participants will be provided with preparatory material in order to better participate in the group exercises.

#### LEARNING OBJECTIVES

At the conclusion of this training course participants wil be able to:

- Apply the basic concepts and principles of signal detection in pharmacovigilance, from simple visualisation and tabulation methods to sophisticated data mining techniques
- Design the signal management process for institutions and companies of various sizes, portfolios and geographical presence, based on the possibilities and limitations of their data and resources
- Understand key messages from the current European and US regulations on signal management, to best manage implications of signals for the future of pharmaceutical companies and products

Participants will complete a knowledge check at the end of the course and will be provided with feedback to ensure learning objectives are attained.

#### **KEY TOPICS**

- Signal detection Theory, methods, data mining
- Signal management Detection, triage, evaluation, further action
- Signal management Regulatory expectations
- Signal management process: Strategy for implementation

## WHO WILL ATTEND

This course is aimed at professionals who work in:

- Pharmacovigilance (including QPPVs)
- Drug safety and patient safety risk management
- Pharmacoepidemiology
- · Information Technology

- · Regulatory Affairs
- Pharmacovigilance consultancies
- Quality and Compliance
- Legal

Course level: Intermediate, for professionals with 2-3 years of experience in Pharmacovigilance, or related functions who are working with Pharmacovigilance around signal management.





#### **Philip Jones**

Medical Director, QPPV inVentiv Health Clinical, UK

## DAY 1

#### **08:00 REGISTRATION**

#### **08:30 WELCOME AND INTRODUCTION**

09:00 SESSION 1

#### SIGNAL DETECTION - THEORY, METHODS, DATA MINING (1)

Signal Detection is a key component of pharmacovigilance science – discovering new effects of medicines in their post-authorisation use. We will present concepts, definitions, data sources, traditional methods of detection, data mining and newest techniques. Participants will exercise the application of best practices in small teams.

#### 10:00 COFFEE BREAK

10:30 SESSION 1 (continued)

SIGNAL DETECTION - THEORY, METHODS, DATA MINING (2)

#### 12:00 LUNCH BREAK

13:00 SESSION 2

# SIGNAL MANAGEMENT - DETECTION, TRIAGE, EVALUATION, FURTHER ACTION (1)

Signal Management is one of the key pharmacovigilance processes; in this session we will review its position relative to other key processes. The different steps and quality requirements in the process will be explained, with the goal of clarifying the need for IT tools and resources for implementation in various contexts. Participants will work on real life examples in small groups.

#### 14:30 COFFEE BREAK

15:00 SESSION 2 (continued)

SIGNAL MANAGEMENT - DETECTION, TRIAGE, EVALUATION, FURTHER ACTION (2)

16:30 SESSION 3

#### **SIGNAL MANAGEMENT - REGULATORY EXPECTATIONS (1)**

We will review the latest regulatory requirements on implementing and maintaining a signal management process. Again, practical exercises will allow participants to understand how they can develop a policy in signal management, and cope with any inspection on this topic.

#### 17:00 NETWORKING RECEPTION

18:00 END OF DAY ONE

# DAY 2

08:00 SESSION 3 (continued)

SIGNAL MANAGEMENT - REGULATORY EXPECTATIONS (2)

09:30 SESSION 4

# SIGNAL MANAGEMENT PROCESS: STRATEGY FOR IMPLEMENTATION (1)

This session reviews how regulators, big companies and small companies should design their signal management process, the expected timelines, resources and outcomes from this process, and how the communication of signals should be performed. We will discuss all of these critical issues during this final session, and look at the typical scenarios the industry and regulators face today.

#### 10:00 COFFEE BREAK

10:30 SESSION 4 (continued)

SIGNAL MANAGEMENT PROCESS: STRATEGY FOR IMPLEMENTATION (2)

12:00 END OF TRAINING COURSE

#### | Plan Your Team's Professional Development

Why not take advantage and train your whole department (or even across different departments!) and benefit from increased:

- Focus
- Flexibility
- Convenience
- Cost Effectiveness

For more information please contact  $\underline{Basel@diaglobal.org}.$ 

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.

# Training Course Venue

#### MERCURE LYON CHARPENNES HOTEL

7 place Charles Hernu 69100 Lyon Villeurbanne, France

Fax: +33 4 78 89 10 14 Email: <u>H1625@accor.com</u>

Tel: +33 4 72 44 46 46

DIA has blocked a limited number of hotel rooms at the rate of EUR 139 from 11 to 14 Nov and EUR 159 from 14 to 16 Nov per standard room per night including breakfast and VAT, excl. EUR 1.65 City-Tax. If you would like to make a booking, please fill in the booking form available on the website and send it per email to <a href="https://doi.org/10.250/jaccor.com">https://doi.org/10.250/jaccor.com</a> with a reference "DIA".

The room rate is available until 15 October 2018 or until the room block is sold-out, whichever comes first.



From train station Lyon Perrache: Take the metro line A, direction L.Bonnevay and exit at station Charpennes (7 stops). From train station Lyon Part Dieu: Take the metro line B to the station Charpennes (2 stops). From the airport Lyon ST EXUPERY: Take the shuttle "Satobus" to the train station TGV/Part Dieu and change there to the metro line B and exit at the station Charpennes (2 stops).



Register 3 individuals from the same company and receive a 50% discount for a 4th! All 4 individuals must register and prepay at the same time without exception. DIA will apply the value of the lowest applicable fee to this discounted 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 only available for the industry rate.

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 to DIA.

For groups of 5 or more individuals, please contact <a href="mailto:Basel@diaglobal.org">Basel@diaglobal.org</a> for a custom group rate.

# **About DIA**

DIA is the global connector in the life sciences product development process. Our association of more than 18,000 members builds productive relationships by bringing together regulators, innovators, and influencers to exchange knowledge and collaborate in an impartial setting. DIA's network creates unparalleled opportunities for exchange of knowledge and has the inter-disciplinary experience to prepare for future developments.

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

# **Continuing Education**

The Faculty of Pharmaceutical Medicine of the Royal Colleges of Physicians of the United Kingdom has accredited this training course with 9.5 CPD credits.

The Swiss Association of Pharmaceutical Professionals (SwAPP) and the Swiss Society for Pharmaceutical Medicine (SGPM) have accredited this training course with 9.5 credits.



# **REGISTRATION FORM**

Signal Management in Pharmacovigilance # 18549 14-15 November 2018 | Mercure Lyon Charpennes Hotel | Lyon, France



#### **REGISTRATION FEES**

Registration fee includes refreshment breaks, lunch on the 1st day of the course and electronic access to training course material.

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	NON-MEMBER
INDUSTRY	€ 1′240.00 □	€ 1′395.00 🗖
ACADEMIA/CHARITABLE/GOVERNMENT/NON-PROFIT (FULL-TIME)	€ 620.00 🗖	€ 775.00 🗖

### All registration fees are subject to applicable French VAT

Please enter your company's French VAT number:

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 non-member fees include a one year membership option. If you registered at one of the non-member rates, you have the opportunity to **become a DIA member** at no additional cost.

To explore membership benefits, please click <u>here</u>. If you want a membership, please indicate your preference below.

☐ I would like to receive a one year complimentary DIA membership at no aditional cost

The DIA Contact Centre Team will be pleased to assist you with your registration from Monday to Friday between 08:00 and 17:00 CET. Tel.:+41 61 225 51 51 Fax: +41 61 225 51 52

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

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 <a href="here">here</a>.

#### **Privacy Policy**

DIA respects the privacy of all of its members and customers. To view our privacy policy, click <u>here</u>. You agree that your personal data will be transferred to DIA in the US

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