





Presented by Management Forum

Medical Device Studies: Regulatory Requirements and Adverse Event Reporting

15-16 September 2025

An essential overview of medical device clinical evaluations, clinical investigations, post-market clinical follow-up requirements and adverse event and vigilance reporting.



Format:

Live online

(1)

CPD:

12 hours for your records

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Certificate of completion

Course overview

This seminar provides an essential overview of medical device clinical evaluations, clinical investigations and PMCF studies with particular emphasis on the adverse event reporting requirements during these studies. The course will concentrate mostly on the new Medical Device Regulations (MDR) with some references to the existing Medical Device Directives.

Benefits of attending

- Gain an insight into medical device clinical studies
- Stay updated of the regulatory changes
- **Learn** the methods of handling adverse events during the study period
- Discuss the new requirements regarding periodic safety update reports (PSURs) and the summary of safety and clinical performance

Who should attend?

- Regulatory affairs specialists
- Quality assurance specialists
- Clinical research associates
- Junior clinical research associates
- Professionals involved in reporting adverse events during pre- and post-market clinical studies

Programme



Clinical evaluation - an overview

- What is a clinical evaluation?
- How do you conduct a clinical evaluation?
- The regulatory requirements pertaining to clinical evaluation

Clinical investigations - an overview

- What is a clinical investigation?
- When are clinical investigations needed?
- The regulatory requirements

PMS and PMCF

- What is PMS?
- What is PMCF?
- When are PMCF studies necessary?

Quiz on clinical evaluation, clinical investigation and PMCF

The competent authority and the Notified Body

- What is their role in the above processes?
- What are the responsibilities of the manufacturer?
- What to communicate and when

Workshop on the new requirements of the MDR



Vigilance workshop

How to define and classify adverse events

- Definitions
- Types of events
- Determining categories

Vigilance reporting

- What is vigilance?
- The requirements for vigilance reporting during medical device studies

The regulatory requirements for monitoring and reporting adverse events during regulatory and PMCF studies

- MEDDEV guidance document
- ISO 14155 harmonised standard
- Responsibilities
- Templates

The MDR - clinical elements

- Focus on changes in the clinical arena
- Chapter 6: clinical evaluation and investigation
- Annex 14: clinical evaluation and post-market clinical follow-up

Course date

15-16 September 2025 Live online

09:30-17:00 **UK (London)** (UTC+01)

Course code 14917

GBP **1,299** 1,499

EUR 1,819 2,099

USD 2,087 2,399

Until 11 Aug

How to book



Online:

ipi.academy/2269

Alternatively contact us to book, or if you have any queries:



Email:

info@ipiacademy.com



Phone:

+44 (0)20 7749 4749

Discounts

- Booking more than one delegate on any one date qualifies for a 30% discount on the second and subsequent places.
- Most events qualify for an early booking discount prior to 6 weeks before the course date. Be sure to check on our website, where the latest discounts will be shown.

Further information

The fee includes all meals and refreshments for the duration of the course (for venue-based courses) and a complete set of course materials (provided electronically). If you have any particular requirements, please advise customer services when booking

Please note

IPI Academy (and our training partners) reserve the right to change the content and timing of the programme, the speakers, the date and venue due to reasons beyond their control. In the unlikely event that the course is cancelled. we will refund the registration fee and disclaim any further liability.

Terms and conditions

The rest of the our terms, the event cancellation policy and the terms and conditions are on our website, please visit ipi.academy/content/terms-and-conditions



Reviews

As the person responsible of safety assessment and reporting in my department (clinical), I was hoping to strengthen my knowledge in this area. Especially when it comes to event assessment and classification. This course helped me achieve that goal. [Speaker] is very knowledgeable on several topics.



Stéphanie Trznadel

Clinical Trial Coordinator SPINEART Sep 16 2024

Understanding what requirements for RA through the cases of Medical Devices studies is very important for me to deal with Korean Medical devices. So this course make me more knowlegable about Europe's MD handling and also distinguish how to adapt on some agenda between Korean and Europe Medical Devices.



Heejun Hong

Student
Dongguk University, Republic of Korea
Oct 3 2022

Fantastic speaker. I truly enjoyed listening to Janette, her ability to relate real life examples helped make sense of the topics.



Melissa Patterson

Associate Manager, Corporate Clinical Compliance Edwards Lifesciences Mar 9 2022

Excellent course, excellent trainer. Found it very valuable and will be taking away knowledge gained to further improve processes and understanding within my department.



Sarah Collins

Pharmacovigilance Manager PharSafer Mar 9 2022

Run this programme in-house for your whole team

Coming to IPI Academy for your in-house training provides an all-inclusive service which gives you access to a wide variety of content, learning platforms and delivery mechanisms as well as your own personal training adviser who will work with you from the initial enquiry through to feedback and follow-up after the programme.

With over 600 trainers, all practitioners and experts across a huge range of fields, we can provide the training you need, where you need it, when you need it, and at a price which suits your budget. Our approach to tailored learning and development consists of designing and delivering the appropriate solution for each client.

For your FREE consultation and to find out more about how we can work with you to solve your training needs, please contact our training advisers:



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