



*Presented by*  
**Management Forum**

# Literature Searching in Drug Safety

**11-12 September 2025**

+ 12-13 January 2026, 19-20 May 2026

This two-day course offers a thorough understanding of literature searching in drug safety, covering regulatory requirements, search criteria, and the interpretation of safety data, while also providing practical assessments to apply knowledge to real-world pharmacovigilance cases.



**Format:**  
Live online



**CPD:**  
12 hours for your  
records



Certificate of  
completion

# Course overview

**This comprehensive two-day course provides an in-depth understanding of literature searching in drug safety, focusing on regulatory requirements, literature search criteria, and the interpretation of safety data from published sources.** Participants will explore best practices for conducting literature searches, identifying relevant safety information, and prioritising articles for review. Practical assessments will enable attendees to apply their knowledge to real-world case studies, ensuring they are equipped with the necessary skills to effectively manage literature searches in pharmacovigilance.

## Benefits of attending

- **Gain** a clear understanding of global and local literature searching regulatory requirements
- **Learn** how assessments help refine literature evaluation and reporting abilities
- **Explore** how to assess safety data to identify potential risks and compliance needs
- **Develop** critical analysis skills for interpreting safety data from literature
- **Discover** how to effectively manage literature searches within pharmacovigilance processes

## Who should attend?

Professionals working within these industries, who want a comprehensive introduction, will benefit from this engaging course:

- Clinical
- Pharmacovigilance
- Regulatory

# Programme

## Day 1

### The regulations concerning literature searchin

- What needs to be searched
- Individual safety data and 'group' data
- The literature search engine requirements – regulatory expectations
- Local literature searching versus global – regulatory expectations

### The literature search criteria

- Regulatory expectations
- Setting up the literature search
- Excipients and literature searching
- Literature search 'hits' and reviews

### What literature articles need to be reviewed

- Individual Case Safety Reports (ICSRs)
- Medical literature monitoring (MLMs)
- Meta-analyses
- Class effect reports
- Clinical studies
- Pharmacoepidemiological studies
- Foreign reports in the literature involving the active ingredient

### Prioritisation of literature articles

- ICSR including abstracts
- Study reports – clinical studies
- Pharmacoepidemiological studies
- Off-label studies

### Literature article reviews

- Literature articles and ICSR
- Literature articles and Periodic Benefit-Risk Evaluation Report (PBRERs)
- Literature articles in signal detection
- Literature articles and 'special situations'

## Day 2

### Practical assessment of literature article and ICSR

- What needs to happen with a literature article report that contains an ICSR
- What needs to happen with the article, timelines and reporting

### Practical assessment of a pre-clinical finding in the literature

- Review a literature article containing information from a pre-clinical study in the literature that contains safety information that is not included in the SPC/Product Monograph and what to do next
- Delegates will be asked what needs to be done with such an article

### Practical assessment of a clinical study

- Delegates will be given a clinical study from the literature conducted by academics for an off-label indication for the company product
- Delegates will be asked what needs to happen with such a report

### Practical assessment of a study involving the company

- Delegates will look at a study involving the company product and asked whether and what should be looked at in the study and where such information should be discussed and reported

### Practical assessment of a pharmacoepidemiological study

- Delegates will be provided with a pharmacoepidemiological study involving the company product
- They will be asked to interpret the information, including what needs to happen next and what if any influence this would have on current labelling or signalling

# Presenter



## **Graeme Ladds**

Graeme is Director of PharSafer and has over 30 years' experience working in the pharmaceutical industry, having started his career at Ashbourne Pharmaceuticals as Head of Drug Safety and Medical Information. Graham has a wealth of experience establishing pharmacovigilance within companies.



# Course dates

## 11-12 September 2025 **Live online**

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

Course code 15681

GBP **1,299** ~~1,499~~

EUR **1,819** ~~2,099~~

USD **2,087** ~~2,399~~

**Until 07 Aug**

## 12-13 January 2026 **Live online**

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

Course code 15682

GBP **1,299** ~~1,499~~

EUR **1,819** ~~2,099~~

USD **2,087** ~~2,399~~

**Until 08 Dec**

## 19-20 May 2026 **Live online**

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

Course code 15683

GBP **1,299** ~~1,499~~

EUR **1,819** ~~2,099~~

USD **2,087** ~~2,399~~

**Until 14 Apr**

## How to book



**Online:**

[ipi.academy/3271](https://ipi.academy/3271)

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



**Email:**

[info@ipiacademy.com](mailto:info@ipiacademy.com)



**Phone:**

[+44 \(0\)20 7749 4749](tel:+442077494749)

## 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

### Fee

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 our terms, the event cancellation policy and the terms and conditions are on our website, please visit [ipi.academy/content/terms-and-conditions](https://ipi.academy/content/terms-and-conditions)

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**IPI**  
Academy

IPI Academy is a training initiative of Falconbury and Management Forum; leading providers of industry training for over 30 years, based in the UK.

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