



*Presented by*  
**Management Forum**

# EU Pharmaceutical Regulations & Strategy

**6-7 October 2025**

The course will provide an up-to-date overview of the current European pharmaceutical regulatory environment including procedures for obtaining marketing authorisation and post-approval obligations. The proposed changes to the EU pharmaceuticals legislation will be discussed in the relevant sections and impact on strategic considerations for obtaining and maintaining marketing authorisations. Procedures for obtaining marketing authorisations in the UK will be discussed.



**Format:**  
Live online



**CPD:**  
12 hours for your records



Certificate of completion

# Course Overview

In the highly regulated field of pharmaceuticals, staying abreast of regulatory affairs is crucial for ensuring compliance, maintaining market authorisations, and effectively managing drug development and post-approval obligations. Navigating the complexities of regulatory processes, particularly in the context of evolving legislation, can be challenging. This is where a comprehensive understanding of pharmaceutical regulatory affairs becomes indispensable.

**Creating and coordinating an effective regulatory strategy is an essential part of the work of a regulatory affairs department and can save valuable time and money. It is therefore important to be aware of potential changes which may impact on strategy.**

This interactive course will provide an overview of the current European pharmaceutical regulatory environment, procedures and obligations and discuss how to interpret and apply the legislation. The proposed EU pharmaceuticals legislation changes will be discussed in the relevant sections.

The programme will cover the legal basis of regulation, development strategies and the importance of pre-submission activities as well a brief overview of the format for presentation of data, the registration procedures for obtaining marketing authorisations and post-authorisation obligations and strategic considerations.

The UK is no longer part of the EU but knowledge of interactions and collaboration with the EU and other regulatory agencies are important for obtaining and maintaining marketing authorisations in the UK. Case study sessions will explore options and strategies for key regulatory activities and provide an opportunity for discussion and the sharing of experiences with our expert trainer and other delegates.

## Benefits of attending

**Understand** the legal basis of the current EU regulatory environment

**Outline** the background to and the proposed EU pharmaceutical legislation changes

**Discuss** development strategy and pre-submission activities

**Review** procedures for applying for a marketing authorisation in the EU/EEA and in the UK

**Discuss** post-authorisation strategic considerations and obligations

## Who should attend?

The course is designed primarily for regulatory affairs personnel, however it will also be of value to those who interact with the regulatory affairs function and would benefit from an understanding of action timelines and information requirements. It will be particularly relevant to all those working in:

- Regulatory affairs
- Project management
- Business planning
- Commercial management
- Manufacturing and QA
- Labelling and artwork
- Medical information
- Clinical
- Pharmacovigilance

# Programme

## Day 1

### EU regulatory environment: legal basis

- Key regulations, directives and guidelines
- Proposed EU pharmaceuticals legislation changes Information sources

### Information sources

#### Case Study One

### Development and Strategy

- Drug discovery
- Scientific advice

### Development process

- Pharmaceutical R&D
- Non-clinical tests
- Clinical studies - Phase I to III

### EU Clinical Trials Regulation

### Types and categories of marketing Authorisations

### Adaptive marketing authorisation procedures

### The Common Technical Document (CTD)

- Overview of Structure and content of a CTD

### Procedures for obtaining a marketing authorisation in the EU and EMA

- The EU centralised procedure

### Other EU centralised procedures

- Referral and arbitration

## Day 2

### Other procedures for obtaining a marketing authorisation in the EU

- Coordination group
- Decentralised procedure (DCP)
- Mutual recognition procedure (MRP)
- National procedures

### Managing product labelling

#### Case Study Two

### Abridged applications and Generics

- Types and Requirements

### Product Life Cycle; Post approval

### Patents and SPCs

### Parallel trade

### Post-authorisation obligations; pharmacovigilance, variations and renewals

### Pharmacovigilance

### Licence variations

- Type I and Type II variations and timelines
- Procedures and timelines

### Extensions

#### Case Study Three

### Renewals

### Sunset clause

### Phase IV Trials

### Classification change

### Generic development

### Strategic factors

### Criteria for successful products

# Presenter



## **Norah Lightowler**

Norah Lightowler is a partner in Lightowler Associates, an independent consultancy offering regulatory advice and support to pharmaceutical companies in or proposing to enter the European market for human pharmaceuticals. They are in their twenty fourth year of successful business. She has wide experience in the pharmaceutical and related nutraceutical, herbal and devices industries as a pharmaceutical assessor with the UK regulatory authority and as associate director of European regulatory affairs with an international pharmaceutical company. She is experienced in organising and presenting courses on European regulatory control systems, including requirements, procedures and strategy.

# Course date

**6-7 October 2025**

**Live online**

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

Course code 15001

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

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

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

**Until 01 Sep**

## How to book



**Online:**

[ipi.academy/2239](https://ipi.academy/2239)

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)

# Reviews



**Overall, it was a wonderful webinar and has helped me further expand my knowledge on the EU/UK regulations. I look forward to participating in future webinars.**



**Iva Martinovic**  
Medical Information Professional  
Teva  
Feb 10 2025



**Very Good. I wanted to get an overview of EU Pharmaceutical Regulations and to understand regulatory strategies to bring new products quickly into the market. The course covered everything I needed and even more.**



**Chitra Saxena**  
Senior Manager Regulatory Affairs  
Lupin Healthcare (UK) Limited  
Feb 10 2025



**Excellent presenter! Truly professional!**



**Anastasios Stylianos Karountzos**  
Regulatory Affairs Officer  
Pharmathen  
Oct 16 2023



**Good overview of the Pharmaceutical Legislation and interesting summary of future changes in the legislation framework. Case studies were a good opportunity to discuss regulatory strategies.**



**Adriana Lopes**  
Regulatory Affairs  
Bluepharma Indústria  
Oct 16 2023

## 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.

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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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**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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