



Presented by
Management Forum

Pharmaceutical Regulatory Affairs in China

26-27 November 2025

A detailed overview of the key areas of pharmaceutical regulatory affairs in China, Hong Kong, Macau and Taiwan.



Format:
Live online



CPD:
12 hours for your records



Certificate of completion

Course overview

This two-day webinar offers a comprehensive review of Pharmaceutical Regulatory Affairs in China. It provides participants with the essential knowledge and understanding of the regulatory requirements for conducting clinical trials in the People's Republic of China (PRC) and for registering pharmaceutical products for the Chinese market.

While the primary focus is on the PRC, the webinar also covers regulatory frameworks in Hong Kong (Special Administrative Region), with brief insights into Macau. Detailed information is provided regarding the registration of pharmaceutical products in both PRC and Taiwan (Republic of China, ROC).

The course also examines pharmaceutical maintenance and other regulatory matters, primarily focusing on innovative products, while including relevant information about generic products.

Please note: This webinar does **not** cover Medical Devices, Immunology drugs, or Vaccines.

All presentation materials will be provided for personal use prior to the webinar. After the event, participants will receive a summary document containing key points, including additional insights shared through the interactive chat functions during the live sessions.

Benefits of attending

- **Gain** access to a wealth of information, which would otherwise be time-consuming to research
- **Exchange** practical experiences with peers and professionals
- **Examine** relevant case studies based on real challenges and opportunities
- **Enhance** knowledge and seek reassurance in solving day-to-day regulatory problems
- **Participate** in frequent Q&A sessions that contribute to enhanced know-how and understanding
- **Receive guidance** on optimising the handling of regulatory situations and challenges

Who should attend

Primarily, this webinar is intended for Pharmaceutical Regulatory Affairs professionals interested in working with China (PR China, including the Hong Kong SAR and Macau SAR), as well as Taiwan (Republic of China), to receive updates on recent developments and to exchange information or seek guidance on current challenges.

Regulatory Affairs professionals who are new to dealing with China and the region, and who require extensive background information and training.

This webinar may also be of interest to other team members from related line functions, including:

- Marketing
- Medicine
- Logistics
- Production, etc., to gain a general overview and introduction

Programme

Day 1

Presentation of participants

Quiz

Drug regulatory systems

Clinical product development

Hong Kong

Macau

Day 2

Product registration strategies

Case studies

Variations

HA Interactions

TAIWAN Republic of China

Recent developments

Q&A and quiz

Presenters



Monica Dressler-Meyer

Mónica Dressler-Meyer is DRA Manager based in Switzerland with many years of regulatory experience. She has spent many years in DRA working with different pharmaceutical companies with responsibility for Asia Pacific and lately also for development activities in other regions. Prior to this, she worked at F. Hoffmann-La Roche in Switzerland where she gained several years' experience in industry basics and pre-clinical research. She has a Degree in Chemistry and Biochemistry from Basel University.



Alan Chalmers

Dr Alan Chalmers is a pharmacist with over 35 industrial experiences mainly in the field of pharmaceutical regulatory affairs. A graduate of Strathclyde University in Glasgow with a B.Sc. in Pharmacy with specialisation in Pharmaceutical Technology, his Ph.D. at Manchester University was in Pharmaceutical Formulation. From 1975-1978 he was Development Officer and Clinical Trials Pharmacist of Allen & Hanburys (part of the then Glaxo group). In 1978 he joined Ciba-Geigy in DRA. Over 20 years were spent with Ciba-Geigy/CIBA/Novartis in all aspects of regulatory affairs including head of a group company DRA in Canada and for many years as Head of Pharma International regulatory affairs.

He has been consultant to the IFPMA, WHO and other international bodies and was Chairman of the Organising Committee of the initial IFPMA Asian Regulatory Conferences in Hong Kong and Singapore and Rapporteur to the more recent conferences in China and Malaysia.

Since 1998 he has been an independent regulatory consultant and is Director of his own consultancy company Pharma International in Switzerland. He has also been director of two UK and Swiss registered pharmaceutical companies with specialised responsibility for international regulatory strategy. More recently as accredited by Swissmedic, Dr. Chalmers is a Qualified Person supporting several Swiss pharmaceutical companies trading internationally with pharmaceuticals and medical devices.

He is published, and his publications include a textbook on *International Pharmaceutical Registration*, *Active Pharmaceutical Ingredients* and as Swiss correspondent to the Regulatory Affairs Journals *Pharma* and *Medtech*. Since 2012 he has been a member of the Editorial Board, *Scrip Regulatory Affairs*.

Course date

26-27 November 2025

Live online

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

Course code 15142

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

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

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

Until 22 Oct

How to book



Online:

ipi.academy/1672

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



Email:

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

Reviews



Very clear, interesting and worth it!



Mélissa Cardenas Trujillo

Regulatory Affairs & Registrations Specialist I
Laboratoires Thea Sas
Apr 29 2024



Valuable Webinar and very good interactions with the speakers.



Celine Aubertin

International Regulatory Affairs Manager
NOOUS FINANCE AU NOM ET POUR LE COMPTE DE
PIERRE FABRE SA
Dec 10 2024



I would highly recommend [...] I appreciate there were more than one speaker to change it up and make the different topics more varied to listen in on. Also very nice how the speakers would supplement each other when responding to questions



Maria Mikkelsen

Regulatory CMC Specialist
Ascendis Pharma
Nov 22 2022



I would highly recommend



Maria Mikkelsen

Regulatory CMC Specialist
Ascendis Pharma
Nov 22 2022

Run this programme in-house for your whole team

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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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