





Presented by Management Forum

Understanding Pharmacovigilance Regulations in APAC

Pharmacovigilance regulations in Asia are dynamic and fastchanging. As the Asia-Pacific region becomes increasingly important for pharmaceutical companies, it's vital to ensure compliance and safety by staying informed of the latest regulatory developments.

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Format: Bespoke training

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CPD: 6 hours for your records (depending on your requirements) ്വ

Certificate of completion

Overview

Pharmacovigilance regulations in Asia are dynamic and

fast-changing. As the Asia-Pacific region becomes increasingly important for pharmaceutical companies, it's vital to ensure compliance and safety by staying informed of the latest regulatory developments.

This two-day pharmacovigilance webinar has been designed to provide a comprehensive guide to compliance in this geographic region. It will provide an ideal opportunity to keep up-to-date with the latest Good Pharmacovigilance Practices (GVP) and post market PV operations in China, and post-market pharmacovigilance regulations in Singapore, Malaysia, the Philippines, and Australia.

Our panel of experts from this region will provide an overview of Good Pharmacovigilance Practices and discuss pharmacovigilance-related requirements. Gain key takeaways to better understand the new GVP regulations and pharmacovigilance compliance in these countries.

Don't miss this opportunity to learn from industry leaders and enhance your knowledge of dynamic and fast-changing regulations in Asia, and the growing market in the Asia-Pacific region.

Benefits of Attending

- **Gain** a comprehensive overview of Pharmacovigilance regulations in China, Singapore, Malaysia, the Philippines, and Australia
- Stay updated with the latest
 Pharmacovigilance regulations in these countries
- Enhance your Pharmacovigilance-related knowledge and skills
- **Understand** the local requirements for RPPV (QPPV), PSMF, RMP, and other related regulations in each of these countries

Who Should Attend

This webinar is intended for anyone involved in and interested in Good Pharmacovigilance Practices (GVP) and the daily practice of pharmacovigilance, including:

- Professionals working in pharmacovigilance departments
- Drug safety specialists
- Adverse reaction monitoring professionals
- R&D professionals
- Regulatory affairs specialists
- Pharmaceutical physicians

Reviews

Overall very well organized. Speakers were very engaging during Q&A session and took time to address every questions. Accomplished what I wanted to learn and more.



Josephine Wong Yunn Shyuan Regulatory Affairs Pharmacist Xepa-Soul Pattison (Malaysia) Sdn. Bhd. Oct 8 2024

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