

AS&T Expert

Job ID

REQ-10045316

Απρ 01, 2025

Malaysia

Περίληψη

The AS &T Expert is responsible for leading the analytical method transfer, method validation, and verification activities, ensuring compliance with cGxP standards for products. Provide technical support to quality control team related to analytics and specifications. Individual contributor role, ensuring the timely implementation of assigned activities.

About the Role

Key Responsibilities:

- Lead analytical method transfer, method validation/verification activities, ensuring full compliance of introduced analytical methods to current standards.
- Act as subject matter expert for analytical and bio-analytical methods for testing of biopharmaceutical molecules (HPLC, UPLC, CIEF, Capillary Electrophoresis, PCR, ELISA, etc)
- Support onsite investigations related to OOX and deviations.
- Customer management and new product introduction to the site.
- Improvements in Analytical Methods and life cycle management.
- Introduction of new technologies for analytical testing.
- APQR, OPV analytics section authoring, and perform assessments.
- Stability Study Design, Protocol, and report preparation. Awareness of ICH requirements.
- Basic knowledge of Statistical data evaluation and trend assessment.
- Regulatory dossier reviews for analytical methods, method validation, stability, and control of materials.
- Third-party testing lab management- Track deliverables and provide technical guidance as per company policy.
- Raw material assessment for new product introduction and pharmacopeia compliance evaluation.

Essential Requirement

- BS degree with 8+ years of pharmaceutical/biotechnology analytics industry experience
- MS degree with 4+ years of pharmaceutical/biotechnology analytics industry experience
- Working experience in Laboratory environment in the biopharmaceutical analytics/QC/ equivalent industry.
- Familiar with major pharmacopeia standards such as USP, EP, JP/JPE, ChP etc
- Broad cGMP experience is required with knowledge and understanding of manufacturing, quality control, and validation requirements and activities
- Must have a working knowledge of health authority and regulatory requirements as well as industry quality management tools, standards, and quality systems.
- Working experience in GMP-regulated industries in the Quality Control analytics department is an

advantage.

- Willing to travel from time to time from Malaysia to Singapore.

Why Novartis: Helping people with disease and their families takes more than innovative science. It takes a community of smart, passionate people like you. Collaborating, supporting and inspiring each other.

Combining to achieve breakthroughs that change patients' lives. Ready to create a brighter future together?

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Τομέας

Operations

Business Unit

Universal Hierarchy Node

Τοποθεσία

Malaysia

Τοποθεσία

Selangor

Company / Legal Entity

MY01 (FCRS = MY001) Novartis Corporation (Malaysia) Sdn. Bhd. (19710100054)

Functional Area

Quality

Job Type

Full time

Employment Type

Regular

Shift Work

No

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List of links present in page

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