QC Specialist II - Analytical

Job ID REQ-10018196 Set 03, 2024 Singapore

Sommario

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About the Role

QC Specialist II - Analytical

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Key Responsibilities:

- Sample storage and management.
- Analytical testing and documentation of API / drug substance / drug product / finished product / Complaints / stability / packaging material samples
- Ensure all activities in compliance with cGxP, incl. data integrity
- Stability (when not centralized)
 - Testing/Sample storage and management
 - Analytical documentation of stability samples to cGxP standards
- Detect and report potential accident, risks and propose solutions

Essential Requirements:

- Preferred: Previous experience working in a laboratory environment in the pharmaceutical industry (quality assurance, production), aseptic technique.
- Administrative activities and GMP and HSE-compliant, efficient production and documentation of standardized tasks in the infrastructure
- Breakthrough Analysis; Being Resilient; Operational Excellence; Continuous Learning; Digital & Tech Savvy
- Laboratory equipment; Quality Control (QC) Testing; Quality Control Sampling; knowledge of TQM and related industry GxP standards and processes; Laboratory Excellence; Quality decision making

Desirable Requirements:

 University degree or equivalent experience in Pharmacy or Chemistry or equivalent + 0-4 years working experience

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Divisione

Operations

Business Unit

Innovative Medicines

Posizione

Singapore

Sito

Tuas South Avenue

Company / Legal Entity

SG12 (FCRS = SG012) Novartis Singapore Pharmaceutical Manufacturing Pte Ltd

Functional Area

Quality

Job Type

Full time

Employment Type

Regular

Shift Work

No

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