

QC Analyst III

Job ID REQ-10048886

4月 24, 2025

Singapore

摘要

-This role utilizes chemistry laboratory skills to test and measure product or materials while ensuring that analysis is performed according to established Standard Operating Procedures (SOPs), Analytical Methods & current Compendia.

About the Role

Key Responsibilities:

- Sample storage and management -Analytical testing/documentation of drug product / finished product / complaints / stability / packaging material samples to GxP standards Stability -Testing/Sample storage and management .
- Analytical documentation of stability samples to GxP standards -Reporting of technical complaints / adverse events / special case scenarios related to Novartis products within 24 hours of receipt -Distribution of marketing samples (where applicable)

Essential Requirement:

- Sound technical & scientific knowledge of pharmaceutical/ chemical.
- Working experience in Laboratory environment in the Pharmaceutical.
- analytics/QC/ equivalent.

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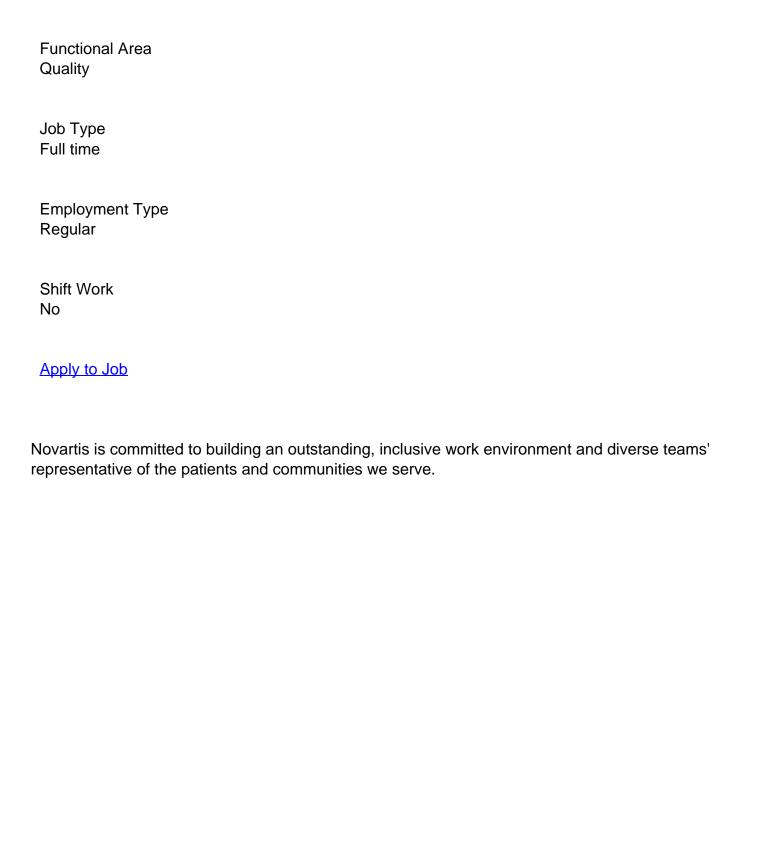
部门 Operations

Business Unit Innovative Medicines

地点 Singapore

站点 Tuas South Avenue

Company / Legal Entity SG12 (FCRS = SG012) Novartis Singapore Pharmaceutical Manufacturing Pte Ltd





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