

QC Analyst II

Job ID REQ-10051659

5月 13, 2025

China

摘要

·此角色利用化学实验室技能测试和测量产品或材料,同时确保分析按照既定的标准操作程序 S(OP、分析方法和当前的编年。

About the Role

Major Accountabilities

- Perform the relevant Analytical testing (product/incoming material testing, stability testing, validation testing, and compliant sample testing, method verification/validation testing, cleaning validation testing, etc.), and provide the testing result in time.
- Execution of assigned tasks in the quality control laboratory in accordance with cGxP regulations. Performance of laboratory specific activities such as maintenance, calibration and qualification of analytical equipment.
- Participate in the establishment, implementation and maintenance of the quality system in lab,

- and support the strength of lab quality system and processes continually.
- Support for investigation related to deviations, OOS/OOE and complaints on analytical topics
- Provides the analytical techniques knowledge and expertise to support QC initiative in quality, compliance and efficiency continuous improvement.
- Comply with all HSE guidelines; Detect and report potential accident, risks and propose solutions
- Responsible for participating in initial training and retraining. Ensure all activities in compliance with cGxP, incl. data integrity

What you'll bring to the role:

- Bachelor Degree in chemistry, pharmacy, biology or equivalent education
- 1-2 years of experience in pharmaceutical industry and/or analytical laboratory in GMP environment desirable
- Language: fluent in Chinese (Oral and written); Basic in English(Oral and written)
- Technical Skills on Laboratory equipment; Quality Control Testing; Quality Control Sampling;
- Collaboration; result-oriented; Breakthrough Analysis; Being Resilient; Operational Excellence; Continuous Learning;

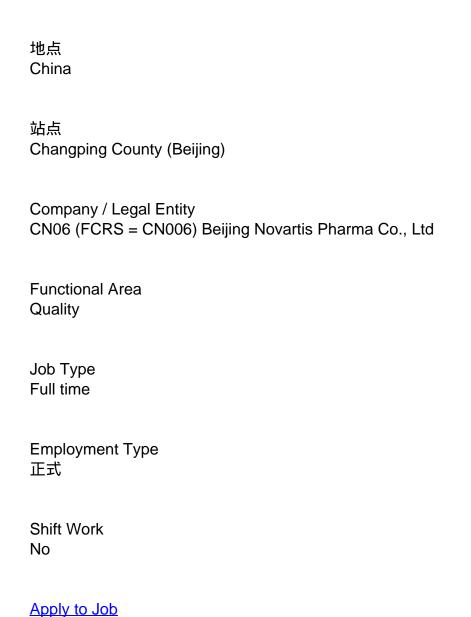
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