

Assoc. Scientist - Technical Development

Job ID
REQ-10080388
Июн. 18, 2026
Индия

Сводка

-Plan and perform scientific experiments (or pilot plant processes) for the preparation and timely delivery of drug substances (DS), drug products (DP), processes and procedures in collaboration within a multifunctional project team coordinated by a Project leader. Contribute to maintenance of lab instruments/infrastructure. -SANDOZ: Plan and perform scientific experiments (or pilot plant processes) for the development and timely delivery of drug products (DP), processes and procedures in collaboration within a multifunctional project team coordinated by a Project leader. Contribute to maintenance of lab instruments/infrastructure. -Support development projects aiming the development of stable, bioequivalent, robust and cost competitive dosage forms -Design and manage experiments/batches for simple/low complexity products under supervision, provide related scientific documentation -Plan and execute experiments in agreement with quality risk management and GDevP /GMP - Assists in the preparation of and reviews of the technological part of dossier

About the Role

• Key Responsibilities

- Plan, perform and document scientific experiments in the laboratory for drug substances (DS) and drug products (DP) in collaboration with multifunctional project teams. Contribute to maintenance of lab instruments/day-to-day operations. Timely execution of project related activities.
- Plan, organize, execute, and document scientific experiments (e.g., analytical method developments/ validations/ transfers/ stability/ release testing, formulation development analytics etc.) according to the agreed timelines and appropriate quality standards.
- Accountable for documentation and submission of raw data in appropriate data system (for e.g., LIMS test activation and results entry).
- Accountable for good documentation practices (GDP) and good laboratory practices (GLP) during execution of laboratory activities.
- Support in evaluation and interpretation of results including investigations on SST failures, OOX/Deviations/Change controls as needed.
- Responsible for implementation and maintenance of lean/efficient/environmentally sustainable practices in the laboratory.
- Proactively communicate key issues and any other critical topics in a timely manner to the manager and/or to any other relevant project team member(s).
- Responsible to meet KQI (Key quality indicators) and KPI (Key performance indicators) for all assigned activities.

Essential Requirements:

- Strong expertise in Chromatography and Dissolution Testing, including method selection and handling
- Proven experience in Analytical Method Development, Validation, and Transfer
- Solid understanding of Regulatory Compliance and GLP standards
- M.Pharm./M.Sc. with up to 2 -5 years of relevant experience, with demonstrated experience in a related domain.

Desirable Requirements:

- Understanding digital tools
- Hands on experience with GLIMS, Chromeleon and ELN

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Development
Business Unit
Development
Место
Индия
Сайт
Hyderabad (Office)
Company / Legal Entity
IN10 (FCRS = IN010) Novartis Healthcare Private Limited
Alternative Location 1
Telangana, Индия
Functional Area
Research & Development
Job Type
Full time
Employment Type
Regular
Shift Work
No

Accessibility and accommodation

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