

Expert Science & Technology

Job ID
REQ-10076531
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Индия

Сводка

As part of this group, you design, plan and/or perform scientific/technical studies. By bridging analytical science to the clinical performance, you will drive the transformation of our molecules into medicines that improve and extend patient's lives. The position is based in the Genome Valley, Hyderabad, within the Technical Research and Development Organization (TRD) of Development

About the Role

Key Responsibilities

- Responsible for performing method feasibilities and validate robust analytical methodologies applied to innovative Oligonucleotides/peptide therapeutics. Strong experience in various chromatography techniques is a pre-requisite. Experience in mass spectrometry applied to biological molecules would be an asset.
- 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 an appropriate data system (for e.g., LIMS test activation and results entry).
- Responsible 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 assigned laboratory related area/activities (e.g., chemical/reagents/consumables/samples/column/ glassware management etc.). 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. Support internal and external audits and ensure no critical findings within the assigned scope.

Minimum Requirements

- Ph.D. in Analytical Chemistry or an equivalent qualification with a minimum of 1–3 years of experience, or M. pharm/M.Sc. with at least 8 years of experience within the pharmaceutical industry, specifically in analytical development. Strong expertise in oligonucleotide/peptide analytics
- Understanding of general regulatory and quality expectations. GMP experience is a must
- Good scientific background, communication skills including presentation and scientific/technical writing.
- Good knowledge of software and computer tools such as Office package, LIMS, chromatography data-evaluation software (e.g. Chromeleon) etc.
- Profound expertise in liquid chromatography separation techniques such as (RP, IEX and HILIC) is a must. Experience in Mass Spectrometry (ranging from mass confirmation to actual quantitative analysis of impurities and sequencing) is an asset
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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?
<https://www.novartis.com/about/strategy/people-and-culture>

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

Accessibility and accommodation

Novartis is committed to working with and providing reasonable accommodation to individuals with disabilities. If, because of a medical condition or disability, you need a reasonable accommodation for any part of the recruitment process, or in order to perform the essential functions of a position, please send an e-mail to diversityandincl.india@novartis.com and let us know the nature of your request and your contact information. Please include the job requisition number in your message.

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