

# **Senior Expert Science & Technology**

Job ID REQ-10005963 Sep 03, 2024 Indien

## Zusammenfassung

400! This is the number of associates in Global Analytical R&D, across 4 countries, working tirelessly on innovative and patient centric medicines. As part of this group, you design, plan and/or perform scientific/technical studies. By bridging the 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 Global Drug Development (GDD).

#### **About the Role**

Your responsibilities will include, but are not limited to:

- Design, plan and perform scientific experiments for projects at different clinical phases of drug substance and drug product with minimal guidance. Well versed with regulatory guidelines, scientific literature, technology transfer and interpretation of the results to draw conclusions in reports.
- Author, review, approve GMP documents (eg: Analytical methods, raw data, SOP's, Qualification reports for analytical instruments.).
- Report and present scientific/technical results internally and contribute to publications, presentations, and patents
- Adhere to Quality metrics, Compliance and Good Documentation Practices following ALCOA+ principles, GLP, OQM, HSE, ISEC and Novartis guidelines.
- Should be a Team player by adding value in collaborating with other teams to support project deliverables within agreed timelines, mentoring new joiners, active participation in project meetings / networks / meetings and contributing to Team goals while meeting individual objectives.
- Ability to perform investigations, guide team members, communicate proactively and clearly to global stakeholders and handle multiple priorities.
- Provide input into CMC documents to support regulatory submission and respond to HA gueries.

#### Role Requirements:

- Desirable knowledge of site language, up to 12 years (for M. Pharma/M.Sc.) & minimum of 6 years (for Ph.D.) experience in Analytical testing & project management in NCE, complex injectables, peptides and late phase method validation.
- Good presentation skills and scientific/technical writing skills.
- · Good communication skills.

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? <a href="https://www.novartis.com/about/strategy/people-and-culture">https://www.novartis.com/about/strategy/people-and-culture</a>

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**Benefits and Rewards:** Read our handbook to learn about all the ways we'll help you thrive personally and professionally: <a href="https://www.novartis.com/careers/benefits-rewards">https://www.novartis.com/careers/benefits-rewards</a>

**Abteilung** 

Development

**Business Unit** 

Innovative Medicines

Ort

Indien

Website

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

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Novartis is committed to building an outstanding, inclusive work environment and diverse teams' representative of the patients and communities we serve.

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REQ-10005963

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