

Expert Science & Technology- Oral Solid Dosage

Job ID
REQ-10074038
мар 16, 2026
Индия

Сводка

Design, plan, perform, interpret, and report results of scientific experiments for the preparation and timely delivery of drug substances (DS), drug products (DP), processes and procedures within global ARD. Lead and manage all project/local network activities, support/coach team members, participate in sub-teams and contribute to overall TRD.

About the Role

Major accountabilities:

- Provide analytical and technical support to PHAD/project team at various stages of product development (eg. CSF, FMI and LCM).
- Design and author analytical documents (e.g., Analytical methods, Stability protocols/reports, Excipient compatibility (EC) protocol/reports; APS protocols/reports, etc.).
- Support Analytical project leader for setting analytical development strategy.
- Support in data interpretation, results compilations and sharing the information with critical observations and proposals to project team.
- Responsible for project related sample handling (e.g., sampling plans, issuance, storage, dis-tribution, reconciliation/destruction of the samples).
- Support planning for assigned project activities. Accountable to meet KQI (Key quality indi-cators) and KPI (Key performance indicators) for all assigned project activities.
- Provide requests for lab activities to the associates and stakeholders.
- Manage project activities including logistics at third parties and external testing laborato-ries.
- Proactively communicate key issues and any other critical topics in a timely manner to the appropriate management level and/or to any other relevant project team member(s).
- Single point of contact for PHAD/project team and other stakeholders (e.g, BioPharm, Mate-rial science and CPP, etc.) for project execution activities.
- Support internal and external audits and ensure no critical findings within the assigned pro-jects.
- Actively contribute to team goals.
- Work according to appropriate SOPs, GMP, GLP, QM, HSE, ISEC & Novartis Guidelines.

Minimum Requirements:

Ph.D in Chemistry/Pharmaceutical sciences with a minimum of 1 years of experience, or M. pharm/M.Sc. with 6 plus years of experience within the pharmaceutical industry, specifically in analytical development.

Work Experience:

- HPLC method development
- Chiral separation

Languages :

- English.

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>

Benefits and Rewards: Learn about all the ways we'll help you thrive personally and professionally.

[Read our handbook \(PDF 30 MB\)](#)

Дивизион

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

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

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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