

Senior Expert Science & Technology-Photochemistry

Job ID 390343BR juin 28, 2024 Chine

Résumé

Design, plan, perform, interpret and report results of scientific experiments for the preparation and timely delivery of drug substances (DS) processes and procedures within a multifunctional project team coordinated by a Project Leader. Manage technical lab/plant activities.

About the Role

Key Responsibilities:

- Independently plan, organize, perform and document scientific experiments /GMP testing /manufacturing plant activities under minimal supervision; handle several activities at a time -Take over responsibility for and utilize special tools /equipment or specialized facilities as an expert; schedule and perform maintenance and qualification of instruments / equipment -Proactively identify conflict situations and contribute to solutions -Work according to appropriate standards for quality, ethics, health, safety, environment protection, and information security; lead initiatives to ensure continuous improvement Documentation of raw data, evaluate and interpret results; propose and actively support the design of next experiments.
- Review and verify raw data generated by others; approval of tests / experiments performed by others Write protocols, scientific reports or lab procedures based on templates or SOPs under minimal
 supervision -For technical development units: Develop new methods or optimize existing
 methods/processes (lab or plant); contribute to development and implementation of new technologies For GMP units: ensure compliance to cGMP -For technology-focused roles: Perform information and
 literature searches under minimal guidance.
- Actively foster knowledge exchange; Train and coach associate scientists, technicians, temporary employees and employees under training / education.
- For project-focused role: Participate in function-specific sub teams and fulfill assigned project tasks and responsibilities under supervision -Uses professional concepts and company's policies and procedures to solve a wide range of difficult problems in imaginative and practical ways.
- Establish innovative solutions for verification and control of critical quality attributes, critical material attributes or critical process parameter in cooperation with other colleagues; Establish control procedures and specifications and review test procedures.

Essential Requirements:

- · Operations Management and Execution.
- Collaborating across boundaries.

· Functional Breadth.

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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Division

Development

Business Unit

Innovative Medicines

Emplacement

Chine

Site

Changshu (Jiangsu Province)

Company / Legal Entity

CN23 (FCRS = CN023) Suzhou Novartis Technical Development Co., Ltd.

Functional Area

Recherche & Développement

Job Type

Full time
Employment Type
Regular
Shift Work
No
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Accessibility and accommodation

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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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