

# **Expert Science & Technology**

Job ID REQ-10010459 Dec 03, 2024 China

# **Summary**

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### **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.

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

## **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

**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/sites/novartis-com/files/novartis-life-handbook.pdf">https://www.novartis.com/sites/novartis-com/files/novartis-life-handbook.pdf</a>

## **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 <a href="mailto:diversityandincl.china@novartis.com">diversityandincl.china@novartis.com</a> 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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Division

Development

**Business Unit** 

Innovative Medicines

Location

China

Site

Changshu (Jiangsu Province)

Company / Legal Entity
CN23 (FCRS = CN023) Suzhou Novartis Technical Development Co., Ltd.
Functional Area
Research & Development
Job Type
Full time

Shift Work

**Employment Type** 

No

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diversityandincl.china@novartis.com

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