

# Clinical Development Medical Manager

Job ID  
REQ-10075503  
апр 12, 2026  
Китай

## Сводка

-Oversees the execution, and interpretation of clinical trials research, data collection activities and clinical operations. Establishes and approves scientific methods for design and implementation of clinical protocols, data collection systems and final reports. Support new and ongoing clinical research and clinical trials and ensure efficient and timely processing of confidentiality agreements and clinical agreements. Monitors adherence to protocols and determines study completion. Manages clinical and regulatory files and maintains clinical inventory intended for distribution to investigational sites. May interact with investigational sites, clinical consultants, Contract Research Organizations and other vendors. Selects, develops and evaluates personnel to ensure the efficient operation of the function.

## About the Role

### Major accountabilities:

- Oversees clinical program(s) across indications, executing medical strategy for development and marketed products in a defined therapeutic area.
- Is responsible for assuring aligned communication with Country/Cluster Clinical Research Associates, Managers and other key stakeholders on the execution and progress of the clinical studies.
- Identify new sites for clinical trials; analyze capability and make recommendation for trial inclusion.
- Facilitate preparation and collection of site level documents; resolve problems as required.
- May execute site initiation and training.
- Implement total site management including monitoring visits, regulatory assessment, drug supply management and resolution of site problems to ensure compliance.
- Track trial execution milestones; identify problems; resolve issues and escalate as appropriate.
- May manage recruitment and execute contingency plans, as needed.
- Complete preparation/generation of study monitoring reports.
- May lead and chair local study team meetings, attend and participate in global clinical trial team meetings -Reporting of technical complaints / adverse events / special case scenarios related to Novartis products within 24 hours of receipt -Distribution of marketing samples (where applicable)

### Key performance indicators:

- Deliver customer satisfaction results for internal and external customers -Delivery of Clinical Trials to quality standards and agreed timelines -Adherence to Novartis policy and guidelines and external regulations.

### Minimum Requirements:

#### Work Experience:

- Managing Crises.
- Functional Breadth.
- Collaborating across boundaries.

#### Skills:

- Clinical Monitoring.
- Clinical Research.
- Clinical Trial Protocol.
- Clinical Trials.
- Decision Making Skills.
- Drug Development.
- Health Sciences.
- Lifesciences.
- Regulatory Compliance.

#### 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  
Китай  
Сайт

Shanghai (Shanghai)  
Company / Legal Entity  
CN14 (FCRS = CN014) China Novartis Institutes for BioMedical Research Co., Ltd.  
Alternative Location 1  
Beijing (Beijing), Китай  
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.china@novartis.com](mailto:diversityandincl.china@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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