

# Clinical Data Scientist - Data Management

Job ID REQ-10049703

Apr 25, 2025

China

## Summary

-Provide timely & professional ongoing Mgmt of Data Mgmt/Coding/CDDRA-Database Development/DAP deliverables and of clinical trial data with respect to cost, quality and timelines for all assigned trials within Clinical Data Mgmt. Ensure consistently high quality data available for analysis and reporting. Develop content and redefine training modules into engaging & interactive applications. Leverage technology to ensure process simplification and training delivery. Follows Good Clinical Practices (GCP), data-handling procedures and guidelines. Participates in the review of clinical research protocols, reports and statistical analysis plans. Drives participation and input within Data Operations (DO) in the delivery of quality data and programs, processes and documentation -Manage data Load, Transfer and conform of Clinical trial data to NCDS compliant standards. The position is a key contributor with Data Provisioning team in ensuring that pharmaceutical drugdevelopment plans in Novartis Global Drug Development are executed efficiently with timely and high quality deliverables.

About the Role

#### Key Responsibilities:

- Provides DM leadership across assigned trial (s) and Acts as the Trial Data Manager where needed -Demonstrates a business understanding of the compound (s) profile to identify and assist in successful application of data Mgmt processes.
- Provides feedback to assure well written protocols and amendments.
- Recognize and resolve protocol issues that may impact database design, data validation and /or analysis/reporting and that do not make the best use of available standards -Performs DM activities for start up of a study, Data Handling plan, Data Review Plan and performing user acceptance testing (UAT) -Manage local lab set up for the Clinical Database as applicable -Leads process and training deliverables within platform or processes.
- Accountable for all aspects of the Process and Training within remit to ensure full compliance to all applicable global regulatory requirements is maintained and business objectives are achieved.
- Accountable for all quality related aspects -Centralizes and aligns DO for audits and inspections.
- Manages and measures Quality -Coordinates exception requests, deviations and corrective
  /preventative action plans -Performs DM hands on activities during the course of the study
  Performs ongoing review of all data generated from the clinical study including Third party
  and local lab data as well as SAE reconciliation where applicable -Responsible and
  accountable to ensure consistency of assigned trials with program level standards across DM
  documentation -Has proven ability to use the tools available to generate listings for data
  review and where necessary provides these to the study teams.
- Generates the study status reports for use at Clinical trial team meetings.
- Supports and assists Junior staff for assigned trials -Provides effective input into DM initiatives and innovations for quality, efficiency and continuous improvement in scientific and operational excellence -Leads /Coordinates synonym review activities and dictionary version upgrade activities at trial /Program level.
- Serves as primary study lead ensuring timely and quality deliverables by establishing and maintaining strong working relationships with study teams, and functional lines.
- Acts as a technical consultant as required.
- Lead DAP activities for assigned /Project level activities for phase I to IV clinical studies in Novartis Global Drug Development.
- Lead independently or participate in improvement initiatives and /or nonclinical projects.
- Reporting of technical complaints / adverse events / special case scenarios related to Novartis products within 24 hours of receipt -Distribution of marketing samples (where applicable)

#### **Essential Requirements:**

- 1-2yrs clinical data management experience
- Bachelor Degree and above
- Fluent English (oral and written). Mandarin Chinese is desirable but not required

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部門 Development
部門 Innovative Medicines
国 China
勤務地 Shanghai (Shanghai)
Company / Legal Entity CN14 (FCRS = CN014) China Novartis Institutes for BioMedical Research Co., Ltd.

Functional Area Research & Development

職種 Full time

雇用形態 Regular

Shift Work

#### Apply to Job.

#### 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.

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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