

Assoc Director, Statistical Programming

Job ID
REQ-10076032
апр 16, 2026
Индия

Сводка

Responsible for all statistical programming/data review reporting & Analytics development aspects of one or more drug development projects or disease area. Direct, oversee and coordinate all activities, deliverables and resources within respective group or disease area. Ensure trials are conducted to a consistently high standard with respect to cost, quality and timelines.

About the Role

Key Responsibilities:

- Lead statistical programming activities for several studies or drive the implementation of data analytics reports -Make decisions and propose strategies at study or project level.
- May act as functional manager for local associates including providing supervision and advice on functional expertise and processes.
- Build and maintain effective working relationship with cross-functional teams, able to summarize and discuss status of deliverables and critical aspects (timelines, scope, resource plan), e.g. as representative in study or project-level team.
- Ensure project-level standardization -Provide and implement programming solutions; ensure knowledge sharing.
- Act as expert in problem-solving aspects.
- Ensure timely and quality development and validation of datasets and outputs for regulatory submissions/interactions, safety reports, publications, post-marketing activities etc -Leads/co-leads novel projects within the team -Generates innovative ideas within own team and /or project team /functional community -Recognizes and leverages innovation opportunities for own team across projects -Mentors and inspires others to solve problems -Reporting of technical complaints / adverse events / special case scenarios related to Novartis products within 24 hours of receipt -Distribution of marketing samples (where applicable)

Minimum Requirements:

- Demonstrates strong proficiency in **SAS** for the analysis and summarization of clinical trial data.
- Has served as a **Trial Programmer** or in a comparable programming role with end-to-end study responsibility.
- Possesses experience in the **development and/or review of critical study documents**, including Protocols, eCRFs, Data Transfer Specifications, SAPs, and mock shells, ensuring consistency with study objectives and regulatory expectations.
- Shows openness to adopting **R and other programming languages**, with a willingness to embrace emerging technologies such as **AI/ML**.
- Exhibits a comprehensive understanding of **CDISC data standards** and their application across clinical studies.
- 10 years' experience in SAS with Graduation min (MSc preferred)

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<https://www.novartis.com/about/strategy/people-and-culture>

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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

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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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