# **U** NOVARTIS

# **Global Clinical Publishing Associate**

Job ID REQ-10006976 Sep 03, 2024 India

### About the Role

Major accountabilities:

• In collaboration with the clinical teams, compile, integrate and publish clinical documents with word processing, electronic publishing, and document management systems in the Novartis Development environment.

• Perform technical quality control (electronic functionality, adherence to internal and external document standards) of published documents.

• Maintain basic knowledge of current electronic publishing standards, regulatory guidelines, and legal requirements.

• Under direct supervision of the immediate manager, acts as the Program Publisher for various programs in clinical development.

Key performance indicators:

- Publish clinical documents (taking into account complexity and size) in accordance with department standards and organization KPIs.
  - Ensure published clinical documents meet current internal and external quality standards for electronic and/or paper HA submissions, including minimizing publishing-related technical QC findings and no rework once finalized.
  - Timeliness of deliverables meet both individual document and overall project timelines.

Minimum Requirements:

Experience with regulatory submission format, including familiarity with submission publishing activities and CTD format criteria.

- Effective interpersonal skills, strong written and oral communication and presentation skills.
- Project management and time management skills to manage multiple ongoing projects simultaneously.

• Familiar with regulatory requirements and HA guidance, including FDA regulations, ICH and EMA guidelines/directives.

- Working knowledge of regulatory affairs.
- Works independently and with minimal supervision.
- Proficiency with computer programs/systems (MS office, etc.) with demonstrated ability to learn new systems quickly.
- Analytical skills and problem solving skills.
- Ability to coordinate and work effectively with cross-functional teams.

Work Experience:

- Cross Cultural Experience.
- Functional Breadth.
- Collaborating across boundaries.
- Operations Management and Execution.
- Project Management.

Skills:

- Clinical Study Reports.
- Data Analysis.
- Documentation Management.
- Lifesciences.
- Operational Excellence.
- Regulatory Compliance.

Languages :

• English.

#### **Role Requirements**

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

Join our Novartis Network: Not the right Novartis role for you? Sign up to our talent community to stay connected and learn about suitable career opportunities as soon as they come up: https://talentnetwork.novartis.com/network

Benefits and Rewards: Read our handbook to learn about all the ways we'll help you thrive personally and professionally: https://www.novartis.com/careers/benefits-rewards

Division Development **Business Unit Innovative Medicines** Location India Site 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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