U NOVARTIS

Associate Clinical Research Medical Director -Immunology, Rheumatology

Job ID REQ-10005168 Mai 22, 2024 USA

Summary

• Accountable for all country clinical/medical aspects associated with Development and prioritized Re-search programs/trials by providing clinical strategic and tactical leadership as the Country Clinical Development representative. May work across several countries.

• Gathers, informs, and acts on clinical/medical/scientific insights for clinical trial concept sheets/protocols, Informed Consent Forms (ICFs) and other relevant clinical documents to optimize clinical trial implementation.

• Drives the identification and involvement of qualified investigators with greatest recruitment potential,

identifies clinical recruitment hurdles and drives clinical recruitment activities to overcome these hurdles.
Accountable for adherence to safety standards and clinical data quality in the country by providing general clinical/medical support for trial related safety findings.

• In close collaboration with other country functions (e.g., clinical trial operations, Medical Affairs, Patient Engagement, and Patient Access) actively contributes to successful allocation, fast clinical trial start-up, timely recruitment, early identification of potential delays, and development and implementation of mitigation plans.

About the Role

Major accountabilities:

- Validates study designs, is accountable for, and makes the final decision on the clinical/medical trial and program feasibility of implementing a clinical trial protocol based on medical/clinical practice and analysis of the competitive environment in the country.
- Actively contributes to scientific/clinical/medical aspects of the start-up phase to ensure fast clinical trial site start-up.
- Provides clinical/medical expertise to clinical trial operations team members and clinical trial sites for Institutional Review Boards (IRB)/ Ethics Committee (EC) interactions.
- Decides on site/Country-specific scientific/clinical/medical content of the Informed Consent Form(ICF) as needed and ensures appropriateness of patient suitable language.
- Provides scientific/clinical/medical expertise during interactions with Country/Cluster external Experts(e.g., Regulatory Authorities, Medical Experts, Advisory Boards, Patient Advocacy Groups, etc.).
- Building disease area expertise, especially for new/rare indications.
- Provides robust indication, compound, and protocol training: To the clinical operations team in the country, especially to the Clinical Research Associates, and other country line functions as needed.
- Supports planning, implementation, and follow-up of scientific/clinical/medical components of Regulatory Authority inspections and internal audits.
- Reviews and resolves Country trial-related scientific/clinical/medical issues/questions. If necessary,

initiates the discussion with the Global Clinical Development team.

- May be involved in reviewing the clinical/medical aspects of clinical trial Serious Adverse Events (SAEs) occurring in the Country and supports the patient safety team, and Global as needed to ensure high guality of clinical/medical information.
- Follows-up with the Investigator for additional clinical/medical information or clarifications for AEs and SAEs and provides clinical/medical expertise for safety amendments. Investigator Notifications (INs). Urgent Safety Measures (USM), etc. as needed.

Job Requirements:

- MD or equivalent medical degree is required, in addition to a proven track record of clinical experience in and scientific contributions to your field of expertise.
- Specialty training in Rheumatology is desired but not an absolute prerequisite.
- Ideally, 3 years of clinical development experience in the pharmaceutical industry or clinical practice
- Working knowledge in the area of Immunology and Inflammation with ability to interpret, discuss and present efficacy & safety data relating to clinical trials.
- Sound understanding of the overall clinical development process, and ICH/GCP principles.
- Ability to manage a study from the scientific/medical/clinical perspective, and a demonstrated capability to problem solve and mediate complex scientific/clinical/medical/operational issues.
- Applies a detailed understanding of the drug in question to provide medical context as it relates to disease processes, populations, and standards of care.
- Ability to assess the feasibility of implementing the protocol based on Country/Cluster medical practice and sound understanding of the overall Clinical Development Plan.
- Demonstrates an understanding of the protocol to evaluate compliance on the part of the Investigator/site staff/study participant and any patient safety issues.
- Demonstrates an understanding of Regulatory requirements and internal policies, procedures, and guidelines pertaining to clinical trials.

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

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Division Development **Business Unit Innovative Medicines** Standort USA Site Remote Position (USA) Company / Legal Entity U014 (FCRS = US014) Novartis Pharmaceuticals Corporation **Functional Area Research & Development** Job Type 2/4

Full time Employment Type Regular Shift Work No <u>Apply to Job</u>

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Major Accountabilities ~ Steigern Sie Ihr wettbewerbsfähiges Umsatzwachstum ~ Identifizierung und Priorisierung von Kunden mit hohem Potenzial durch Datenanalyse (HCPs und Stakeholder), die Verschreibungsentscheidungen beeinflussen ~ Steigern Sie die Vertriebsleistung durch die geschickte Orchestrierung positiver Kundenerlebnisse ~ Engagieren und Beziehungen aufbauen ~ Führen Sie wertorientierte Gespräche (persönlich und virtuell), um kritische Kundenherausforderungen, Entscheidungstreiber, Schwachstellen und Chancen zu verstehen ~ Personalisieren und orchestrieren Sie Customer Engagement Journeys für HCPs,

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