

Clinical Research Medical Advisor

Job ID
REQ-10056559
Jul 02, 2025
Australia

Summary

Location: Sydney, Australia #LI-Hybrid

Novartis is unable to offer relocation support for this role: please only apply if this location is accessible for you.

About the Role:

As the Clinical Research Medical Advisor (CRMA), your responsibility lies within the development of Global Clinical Trials – this includes medical oversight for all trials, portfolio and/or protocol medical feasibility, scientific engagement of investigators, protocol and TA training for internal and external stakeholders, medical issue or question management, safety review, strategic input in pre-launch planning.

You will drive compliance across all aspects of clinical trials and CRMA related activities. It will be critical to ensure good communication and stakeholder management cross-functionally within the local country organisation as well as between global and regional teams.

About the Role

Key Responsibilities:

- Medical oversight of clinical trials across all stages and contribute to operational trial deliverables, according to timelines, quality/compliance, and performance standards.
- Drive portfolio/trial medical feasibility within the Global Development framework and provide country clinical strategic guidance and proposals in collaboration with Study and Site Operations Team and Medical Affairs Team.
- Identify and propose new sites for clinical trials, analyse capability, assess patient pool and country treatment landscape, and make recommendations for potential trial inclusion.
- Provide robust indication and protocol training to CRAs, CSMs, RSMs and other functions in the country as needed.
- Responsible for medical related education, implementation and compliance to protocol, standards (SOPs) and best practices for clinical development within assigned clinical trial(s) and within clinical program(s), including sharing lessons learned.
- Provide medical expertise to clinical/operational activities for patient eligibility, medical question-management, safety, amendments, etc.
- Collaborate cross-functionally for the early product launch planning process to ensure Global Development trials conducted are aligned with the local country strategy.
- Support medical/clinical team discussions with local regulatory interactions as needed.

Essential Requirements:

- Medical Degree (MD, MBBS).
- Proven experience in medical practice or pharmaceutical industry experience with a background in clinical trials/medical affairs/life sciences/research in all aspects of drug development including clinical research, GCP, and local regulatory requirements.
- Experience in Haematology and Oncology clinical trials is valuable.
- Demonstrated experience in managing projects, feasibility conduct and the execution of strategic plans from a medical perspective.
- Outstanding internal and external stakeholder engagement experience.
- Location is based in Sydney, with flexible working options.

Commitment to Diversity and Inclusion / EEO paragraph:

Novartis is committed to building an outstanding, inclusive work environment and diverse teams representative of the patients and communities we serve.

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?
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Division

Development

Business Unit

Universal Hierarchy Node

Location

Australia

Site

New South Wales (NSW)

Company / Legal Entity

AU04 (FCRS = AU004) AU Pharma Pty Ltd

Functional Area

Research & Development

Job Type

Full time

Employment Type

Regular

Shift Work

No

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