

# **Clinical Research Medical Advisor**

Job ID REQ-10018683 Ago 20, 2024 Países Bajos

#### Resumen

As a Clinical Research Medical Advisor, you will play a key role in providing clinical, strategic and tactical leadership to support Global Drug Development (GDD) trials and clinical development plans with a medical / clinical accountability and direct impact on the efficiency of clinical trial conduct.

#### **About the Role**

# As a CRMA your responsibilities include, but are not limited to:

- •Closely collaborates with Study & Site Operations (SSO) to ensure fast clinical trial start up,recruitment according to planned timelines, early identification of potential delays and robust recruitment mitigation plans. Co own start up phase and the recruitment plan for the development clinical trials with the local SSO organization.
- •Provide clinical development and indication expertise and drives, together with the local SSO, the execution of clinical trials with high quality and within planned timelines; provide indication and protocol training's to investigational sites as well as SS O colleagues.
- •Assess the clinical feasibility of implementing a clinical trial protocol based on regional/local medical practice using physician interviews, local databases (RWE, payer data, patient association feed back, etc.) and analysis of the competitive environment.
- •Cooperates with local functions such as e.g. Medical Affairs, Patient access to identify and involve qualified investigators with recruitment potential and relevant key experts for clinical development in order to exploit the value of the assigned project(s) in the context of the investigational product(s).
- •Review and resolution of local medical issues / questions if necessary, support the discussion of issues to global teams.

#### **Requirements:**

- Scientific degree ideally MD
- •experience in the pharmaceutical industry with at least 3 years' experience in clinical development or trial monitoring across different indications / therapeutic areas.
- Fluent English
- •Ability to manage a study from the medical / clinical perspective, and a demonstrated capability to problem solve and mediate complex clinical / medical / operational issues.
- •Demonstrate an understanding of regulatory requirements and policies, procedures, and guidelines pertaining to clinical trials.
- •Track record of delivering complex global clinical projects in quality and time.
- •Excellent communication and interpersonal skills, with ability to build rapport and trust with diverse stakeholders.
- •This position will require local and international travels; up to 30% of working time

#### Why Novartis:

Our purpose is to reimagine medicine to improve and extend people's lives and our vision is to become the most valued and trusted medicines company in the world. How can we achieve this?

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**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? <a href="https://www.novartis.com/about/strategy/people-and-culture">https://www.novartis.com/about/strategy/people-and-culture</a>

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División

International

**Business Unit** 

**Innovative Medicines** 

Ubicación

Países Bajos

Sitio

Amsterdam

Company / Legal Entity

NL08 (FCRS = NL008) Novartis Pharma NL

**Functional Area** 

Research & Development

Job Type

Full time

**Employment Type** 

Regular

Shift Work

No

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