

# Senior Clinical Development Medical Director, Oncology

Job ID 393918BR

8月 02, 2024

**USA** 

# 摘要

Onsite

Location: East Hanover, New Jersey

Hybrid #LI-Hybrid

#### About the role:

Novartis is deeply committed to transforming the lives of people living with solid tumors, blood cancers and serious or life-threatening blood disorders. We believe that anyone living with these conditions has the right to a life free from pain, free from symptoms and free from disease - this is our vision for the future.

As the Senior Clinical Development Medical Director (CDMD), you will lead the strategic planning and management of the assigned clinical program from an end-to-end clinical development perspective. As Sr CDMD, you will have oversight of the clinical development for the assigned programs and drive execution of the clinical development plan. You will enable an empowered organization, which can navigate in a matrix environment and adjust quickly to business needs.

In Clinical Development Oncology, our aim is to design innovative, patient friendly clinical development plans to rapidly bring outstanding treatments to patients, caregivers and healthcare systems. We are striving to develop treatments for Lung, Breast & Prostate Cancers, MDS & AML, CML and sickle-cell disease, and are pushing the boundaries of innovation with CAR-T and Radioligand therapies.

# About the Role

# Your Key Responsibilities:

- Providing clinical leadership and strategic medical input for all clinical deliverables in the assigned project or section of a clinical program
- Leading development of clinical sections of trial and program level regulatory documents
- Driving execution of the assigned clinical program and/or clinical trial in partnership with global line functions, assigned Global Trial Directors (GTDs), and regional/country medical associates, where applicable
- Support the Global Program Clinical Head (GPCH) in ensuring overall safety of the molecule for the assigned section, and may act as a core member of the Safety Management Team (SMT), supporting overall program safety reporting in collaboration with Patient Safety colleagues
- Supporting the Clinical Development Head (CDH) by providing medical input into Clinical Development Plan (CDP), Integrated Development Plan (IDP) and Clinical Trial Protocol (CTP) reviews, and contributing to/driving development of disease clinical standards for new disease areas
- As a medical expert, supporting the GPCH or CDH in interactions with external and internal stakeholders and decision boards

The ideal location for this role is East Hanover, NJ, but remote work may be possible (there may be restrictions based on legal entity). Please note that this role would not provide relocation as a result. If the associate is remote, all home office expenses and travel/lodging to the East Hanover or corporate site for periodic live meetings will be at the employee's expense. The expectation of working hours and travel (domestic and/or international) will be defined by the hiring manager.

Video Link <a href="https://www.youtube.com/watch?v=ggbnzRY9z8w">https://www.youtube.com/watch?v=ggbnzRY9z8w</a>

### Role Requirements:

# Essential Requirements:

- MD or equivalent medical degree is required in addition to advanced knowledge and clinical training in medical/scientific area; Clinical practice experience: 4 years (including residency) preferred.
- Minimum of 7 years of experience in clinical research or drug development.
- Experience in an academic or industry environment spanning clinical activities in Phases I-4 required.
- 2 years of contribution to and accomplishment in all aspects of conducting clinical trials (e.g.,

- planning, executing, reporting and publishing) in a global/matrix environment in pharmaceutical industry required.
- Working knowledge of Oncology is required, with proven ability to interpret, discuss and present efficacy and safety data relating to clinical trials.
- Demonstrated ability to establish effective scientific partnerships with key stakeholders.
- Working knowledge of GCP, clinical trial design, statistics, and regulatory and clinical development processes.
- Previous global people management experience is preferred, though this may include management in a matrix environment.

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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### Accessibility & Reasonable Accommodations

The Novartis Group of Companies are 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 application process, or to perform the essential functions of a position, please send an e-mail to <u>us.reasonableaccommodations@novartis.com</u> or call +1(877)395-2339 and let us know the nature of your request and your contact information. Please include the job requisition number in your message.

部门 Development

Business Unit Innovative Medicines

地点 USA

站点 East Hanover

Company / Legal Entity U014 (FCRS = US014) Novartis Pharmaceuticals Corporation

Functional Area Research & Development Job Type Full time

Employment Type Regular

Shift Work No

# Apply to Job

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