

## Senior Clinical Development Director

Job ID REQ-10051511

6月 14, 2025

USA

## 摘要

The Sr Clinical Development Director (CDD) in the Cardio Development Unit is responsible for leading the strategic planning and management of the assigned clinical program(s) from an end-toend clinical development perspective. As a CDD in the TA, you will have oversight of the clinical development for the assigned programs and drive the execution of the clinical development plan. In addition, you will enable an empowered organization, which can navigate in a matrix environment and adjust quickly to business needs.

## About the Role

Major accountabilities:

• Providing clinical leadership and strategic input for all clinical deliverables in the assigned project or section of a clinical program. Clinical deliverables may include clinical sections of individual protocols or sub studies consistent with the Integrated Development Plans (IDP),

clinical data review, program specific standards, clinical components of regulatory documents/registration dossiers, and publications

- · Leading development of clinical sections of trial and program level regulatory documents
- Driving execution of the section of the clinical program in partnership with global line functions, assigned Global Trial Directors (GTDs), and regional/country medical associates, if applicable
- Overseeing/conducting ongoing medical and scientific review of clinical trial data with Clinical Scientific Expert(s) with appropriate oversight from Medical Lead
- Supporting (Sr.) GPCH in ensuring overall safety of the molecule for the assigned section and may be a core member of the Safety Management Team, supporting overall program safety reporting in collaboration with Patient Safety
- As a clinical expert, supporting the (Sr.) GPCH or CDH in interactions with external and internal stakeholders and decision boards
- Contributing to scientific training of relevant Novartis stakeholders on the disease area and compound/molecule. May serve as speaker for franchise medical/scientific training and may be the Program Manager of other associates

Work Experience:

- Advanced degree in life sciences/healthcare (or clinically relevant degree) is required. PharmD, or PhD strongly preferred
- 10 years of involvement in clinical research, global drug development in an academic or industry environment spanning clinical activities in Phases I through IV. 5 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. Experience in late phase clinical development strongly preferred
- Solid scientific writing skills
- Experience with regulatory submissions (IND, NDA/BLA, CTA/MAA) preferred
- Solid and advanced scientific acumen and ability to analyze and interpret scientific literature and data. Strong affinity with data, data quality and analysis.
- Preferred knowledge and/or experience of assigned therapeutic area
- Demonstrated ability to establish strong scientific partnership with key internal and external stakeholders
- 3 years people management experience required; this may include management in a matrix environment\*

The salary for this position is expected to range between \$204,400 and \$379,600 per year.

The final salary offered is determined based on factors like, but not limited to, relevant skills and experience, and upon joining Novartis will be reviewed periodically. Novartis may change the published salary range based on company and market factors.

Your compensation will include a performance-based cash incentive and, depending on the level of the role, eligibility to be considered for annual equity awards.

US-based eligible employees will receive a comprehensive benefits package that includes health, life

and disability benefits, a 401(k) with company contribution and match, and a variety of other benefits. In addition, employees are eligible for a generous time off package including vacation, personal days, holidays and other leaves.

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? <u>https://www.novartis.com/about/strategy/people-and-culture</u>

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: <a href="https://talentnetwork.novartis.com/network">https://talentnetwork.novartis.com/network</a>

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

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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.

Business Unit Universal Hierarchy Node

地点 USA

状态 New Jersey

站点 East Hanover

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

Alternative Location 1 Cambridge (Massachusetts), Massachusetts, USA

Functional Area Research & Development

Job Type Full time

Employment Type Regular

Shift Work No

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