

Study Monitor-Principal Scientist, Translational Medicine/Preclinical Safety (PCS) Remote Position

Job ID
REQ-10080327
Июн. 18, 2026
США

Сводка

LI#-Remote
Internal Title: Principal Scientist I/II

The Preclinical Safety (PCS) department within the Biomedical Research (BR) - Translational Medicine Unit provides non-clinical safety strategy of products in - discovery, -development and -market, globally, with state-of-the-art regulatory compliance.

As a Scientific Study monitor, you will join our PCS team to oversee non-clinical research activities for multiple projects across multiple disease areas for in-vivo toxicity and /or in-vitro screening toxicity studies conducted at our CRO partner sites as per the internal strategy and international standards, acting as the primary scientific contact for the Study Director.

About the Role

Key Responsibilities:

- Formulates and leads/co-leads novel projects with team or enables matrix collaboration on project/technology solutions to achieve creative results for impact on BR goals. Generates innovative ideas within own team and/or project team/functional community to meet new technical requirements and/or answer project key scientific/technical/development questions. Establishes target dates and priorities to enable data-driven advancements in project teams, within own team, and with collaborators, or within functional community.
- The Study Monitor is appointed to each outsourced preclinical study based on relevant technical expertise, designated disease area and/or scientific background knowledge and acts as the primary scientific contact for the Study Director at the Contract Research Organization (CRO).
- The Study Monitor is responsible for overseeing the progress of the study and for ensuring that the study is conducted, recorded and reported according to the study protocol. The Study Monitor should ensure that the study is compliant with the appropriate GLP regulations, Novartis animal welfare policies, CRO in-house standard operating procedures, Novartis expert recommendations (where feasible) and all relevant international regulatory guidelines/regulations.
- Resolution of study related issues, liaisons with internal experts and informing the appropriate people in a timely manner is pivotal to the performance of this role. The study phases and sample delivery timelines should be strategically overviewed and tracked to ensure that internal contributor reports are delivered to the CRO on time and that the CRO meets its agreed main reporting timelines.
- Communication skill is critical to this role in forming strong working relationship with other Target team members.
- Works closely with the PCS-Operations and PCS Project Team Member (PTM) to formulate a project outsourcing strategy.
- Has a working knowledge of HA regulations (Swiss medic, OECD, FDA) to support conduct of GLP compliant toxicology studies.
- May be PCS part-time PTM

Essential Requirements:

- PhD or MVSc/MS/M.Pharm with 7+ years of experiences in drug discovery and/or development, preferably as Study Director or Study Monitor in the early preclinical screening and GLP studies
- In-depth knowledge of toxicology assays in early development, Safety pharmacology and genotoxicity
- Proficient with full range of techniques used in job and core areas. Working knowledge of tools and processes used in drug design and development.
- Excellent communicators, strong team players and have a high level of logistical/planning ability.
- Registration and certification with one of the International Toxicology reg

This is a dual posting. The final level & title of the offer role would be determined by the hiring team based on the skills, experience & capabilities required to perform the role at the level the role has been offered.

The salary for this position is expected to range between:

Principal Scientist I: \$119,700 and \$222,300 per year.

Principal Scientist II: \$126,000 and \$234,00 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. To learn more about the culture, rewards and benefits we offer our people click [here](#).

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>

Benefits and Rewards: Learn about all the ways we'll help you thrive personally and professionally.

[Read our handbook \(PDF 30 MB\)](#)

EEO Statement:

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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 us.reasonableaccommodations@novartis.com 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.

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Functional Area
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