U NOVARTIS

Associate Clinical Development Medical Director

Job ID REQ-10030826 Nov 21, 2024 Ireland

About the Role

Major accountabilities:

Your responsibilities as a Nuclear Medicine expert will include:

 Providing clinical leadership and strategic medical input for all clinical results in the assigned project or section of a clinical program

Leading development of RLI related clinical sections of trial and program level
regulatory documents

• Driving execution of the program and/or clinical trial in partnership with global line functions, assigned Global Trial Directors (GTDs), and regional/country medical associates, where applicable

• Supporting (Senior) 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 Nuclear Medicine physician specialist, supporting the (Sr.) GPCH or CDH in interactions with external and internal partners and decision boards

Contribute to the publication strategy of RLI/RLT compounds from the scientific standpoint

• May work with BR (Biomedical Research)/ Translational Medical Sciences) to drive transition of pre-PoC (Proof of Concept) projects to DDP (Development Decision Point) and with BD&L (Business Development & Licensing) including target identification and due diligences together with other medical matters, as needed.

Role Requirements :

Nuclear medicine Physician/Medical Doctor

• Sophisticated knowledge and clinical training in oncology PET; Clinical practice

experience \geq 5 years preferred.

- Experience in Clinical Trials with a PET component
- Experience with Radioligand therapy

• A consistent track record to interpret, discuss and present data relating to clinical trial(s) with a Nuclear Medicine component

Demonstrated ability to establish effective scientific partnerships with key partners

• Solid understanding of GCP, clinical trial design, statistics, regulatory and clinical development processes

Role Requirements

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

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: https://talentnetwork.novartis.com/network

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>

Division Development **Business Unit Innovative Medicines** Location Ireland Site Dublin (NOCC) Company / Legal Entity IE02 (FCRS = IE002) Novartis Ireland Ltd Alternative Location 1 United Kingdom Alternative Location 2 Spain **Functional Area Research & Development** Job Type Full time **Employment Type** Regular Shift Work

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