

RLT Medical Launch Liaison

Job ID REQ-10002981 Apr 22, 2024 China

Summary

About the role:

- RLT MLL serves as an ambassador and an advocate for AAA science within the healthcare community, and works directly with Nuclear medicine/oncology healthcare professionals across accounts, educates targeted radioligand treatment centers about appropriate setup, process, handling, administration and infrastructure, and as well as to discuss and provide scientific and therapeutic information on appropriate radiopharmaceutical licensing for Lu177, Ga 68 and other relevant radio linkers
- In line with overall product strategy, the RLT MLL is responsible for supporting regional strategy, implementation and execution of Medical Affairs plans for radioligand therapy and radioligand imaging, providing scientific information, helping design and organize clinical studies, building educational dialogue with KOLs and regulatory stakeholders
- Provide regional RLT leadership to accelerate the registration for new pipeline

About the Role

Key Responsibilities

- •Be a strategic internal partner, gather and leverage insights for an impactful contribution to Patient Journey mapping, launch excellence roadmap, integrated evidence generation plans, integrated product strategy and subsequently the medical strategy.
- •Utilize knowledge of assigned therapeutic area and Novartis compounds to serve as the Medical, Clinical and Scientific expert to field matrix colleagues
- •Ensure appropriate identification and mapping of external stakeholders, aligned to the medical strategy, and in collaboration with other Novartis colleagues. Able to create personalized, flexible engagement strategies and plans, leveraging multiple channels and tailored content to meet the changing needs of external stakeholders
- •Respond to unsolicited requests for information from stakeholders by sharing appropriate data regarding marketed and pipeline compounds in a timely, compliant, and stakeholder-focused manner.
- •Manage administrative responsibilities in a timely manner (customer relationship management tool, compliance training and other modules, expense reporting, etc.).
- •Promote and adhere to Ethics and Compliance Professional Practices Policy (P3).
- •Reporting of technical complaints / adverse events / special case scenarios related to Novartis products within 24 hours of receipt. Distribution of marketing samples

Essential Requirements:

- •MD, PharmD., pharmacist, PhD, Master above degree in medical or other life sciences
- Ability of Chinese and English
- •Medical Science and Disease Area Knowledge

- Strategic& Enterprise Mindset
- Stakeholder Engagement& Experience
- •Healthcare systems& partnerships
- Joint Value Creation Collaboration

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: Read our handbook to learn about all the ways we'll help you thrive personally and professionally: https://www.novartis.com/sites/novartis com/files/novartis-life-handbook.pdf

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Novartis is committed to building an outstanding, inclusive work environment and diverse teams' representative of the patients and communities we serve.

Division

International

Business Unit

Innovative Medicines

Standort

China

Site

Chengdu (Sichuan Province)

Company / Legal Entity

CN06 (FCRS = CN006) Beijing Novartis Pharma Co., Ltd

Functional Area

Research & Development
Job Type
Full time
Employment Type
Regular
Shift Work
No
Apply to Job

Accessibility and accommodation

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EEO Statement:

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