

# **Medical Director, Renal**

Job ID REQ-10029229 Ene 09, 2025 Reino Unido

#### Resumen

- ~ Entwickelt und implementiert strategische und operative TAs Global Medical Affairs-Programme mit Schwerpunkt auf innovativer Evidenz und/oder Markteinführungsbereitschaft und/oder Lösungen für die Zeit nach der Markteinführung, einschließlich der Planung medizinischer Angelegenheiten und der Umsetzung der medizinischen/wissenschaftlichen Engagement-Strategie, die sich mit den Bedürfnissen strategischer medizinischer Aktivitäten vor der Markteinführung und Markteinführung befasst und diese erfüllt, um den Bedarf an Patienten, Kliniken, Zugang und Wert für Gesundheitssysteme zu gewährleisten
- ~ Bietet Fachwissen in der Entwicklung und Ausführung der übergreifenden Strategien, liefert Inputs während des Designs und entlang der End-to-End-Ausführung von Programmen
- ~Entwickelt und führt den Integrated Evidence Plan (IEP)/funktionsspezifische Programme aus, um das Wertversprechen für das priorisierte Markteinführungsportfolio und die Wirkung unserer Medikamente zu maximieren.

## **About the Role**

#### Major Accountabilites:

- Lead development and execution of medical affairs strategy for Renal priority programs including transformative tactics such as: research/population health, innovative partnerships and integrated evidence plans
- Co-develop plans for evidence generation, MSL / Field Medical Affairs strategy, medical education programs, scientific publication planning and Medical Expert network development with TAs
- Co-own the development and implementation of innovative education and scientific communication plans for external stakeholders
- Financial tracking to ensure timely and cost-effective development & execution of medical activities
- Prepare SRC submissions for TA assets within remit
- Partner with Development, S&G, US and International cross-functions to shape portfolio early and diversify evidence to achieve broad access at launch and to enhance impact on clinical practice for priority programs
- Represent GMA around prioritized portfolio with internal and external audiences, in collaboration with TAs including the investment, medical and regulatory communities, as well as pharmaceutical or biotechnology industry collaborators/partners
- Represent "the voice of the patient" internally and evaluate factors relevant to a patient's informed decision making
- Provide direction and input into the development and implementation of successful reimbursement and market-access strategies
- -Provide proactive input to Development on potential new therapeutic indications, to enrich Registration Programs and to consider new therapeutic opportunities.
- Ensure that Patient Access programs are supported for all brands within the GMA and delivered with full

#### compliance

- Ensures GMA activities are designed and executed in compliance with company policy guidelines and highest medical quality standards
- Provide proactive medical input to asset lifecycle management to consider new therapeutic opportunities
- Ensure that Patient Access programs are supported for all brands within International Medical Affairs and delivered with full compliance

# Requirements:

#### Must have:

- MD (Preferred) or PhD/PharmD in Health Sciences. Specialist Degree or specialist qualification related to discipline for which you will be responsible is an advantage.
- 5+ years in Pharmaceutical Industry experience in Medical Affairs and/or Clinical Development
- Critical thinker and with ability to navigate uncertainty without major supervision
- Fluent oral and written English; Other relevant languages are an advantage.
- Strategic mindset and able to establish credibility and influence across a range of diverse stakeholders in a matrix organization to drive change
- Ability to truly collaborate across functions and markets: serve-partner-co-create
- Able to navigate in an environment of shared outcomes and cross-business accountabilities
- Deep understanding of health care systems and key external stakeholders
- Strong track record of delivery focus for time and quality in medical affairs projects
- Successful development and implementation of innovative programs and processes
- Understands unmet medical needs, generates the right evidence to fulfil them, uses innovative, multichannel communication formats for effective evidence dissemination
- Credibility as peer expert with external stakeholders
- · Agile mindset & ability to lead in an agile organization across Disease Areas
- Firm working knowledge of GCP, scientific and clinical methodology, protocol designs, management and regulatory requirements for clinical studies designated for review by regulatory authorities.

#### Preferred

- Highly preferred: Renal experience particularly IgA Nephropathy, significant medical affairs early asset lifecycle, pre-launch and launch experience in Global organizations
- Rare disease experience
- Experience in developing and executing "Best in Class" processes at scale
- Clinical trial research experience conducted in a pharmaceutical or equivalent academic environment in TA
  of interest is strongly desired

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División

Development

**Business Unit** 

Innovative Medicines

Ubicación

Reino Unido

Sitio

London (The Westworks)

Company / Legal Entity

GB16 (FCRS = GB016) Novartis Pharmaceuticals UK Ltd.

Alternative Location 1

Barcelona Gran Vía, España

Functional Area

Research & Development

Job Type

Full time

**Employment Type** 

Regulär

Shift Work

No

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