

# Medical Head Solid Tumors

Job ID REQ-10034455

1月 10, 2025

Italy

## 摘要

Sviluppa e implementa programmi strategici e operativi di TA Global Medical Affairs, con particolare attenzione alle evidenze innovative e/o alla prontezza al lancio e/o alle soluzioni post-commercializzazione, compresa la pianificazione degli affari medici e l'esecuzione della strategia di coinvolgimento medico/scientifico che affronta e fornisce le esigenze strategiche delle attivit à mediche pre-lancio e lancio per i pazienti, cliniche, l'accesso e il valore ai sistemi sanitari Fornisce competenze nello sviluppo e nell'esecuzione delle strategie generali, fornendo input durante la progettazione e lungo l'esecuzione end-to-end dei programmi Sviluppa ed esegue l'Integrated Evidence Plan (IEP)/programmi funzionali specifici per massimizzare la proposta di valore per il portafoglio di lanci prioritari e l'impatto dei nostri farmaci.

About the Role

Key Responsibilities:

Your responsibilities include, but are not limited to:

- Develop and implement the overall strategic direction for the Solid Tumors medical department in alignment with the goals and objectives of the broader oncology area and of the organization
- Continuously monitor industry trends, research advancements, and regulatory changes to adapt the strategy and ensure compliance with best practices
- Provides medical leadership, ensuring cross-functional alignment with the other teams inside and outside Medical Department to establish and execute strategic initiatives to improve patient outcomes and ensure the provision of cutting-edge Solid Tumors services.
- Supervise effective clinical development of products through accountability of the local medical plans for the Solid Tumors TA.
- Foster a culture of continuous learning, research, and innovation to drive advancements in Solid Tumors treatment options and care delivery.
- Lead scientific interactions with Top Medical Experts and Scientific Society of the Solid Tumors Area.
- Responsible for medical approval as per Doing Business Ethical (materials, events, grants as examples).
- Drive collaboration of the local medical team with Local, Regional and Global Teams in local Clinical Development activities and in Global and International Med Affairs activities
- Oversee together with Patient Safety local clinical trial adverse event reporting as well as clinical input and oversight into adherence to GCP together with SSO.
- Present highest ethical standards and contribute proactively to a credible reputation for Novartis CPO in the local Health Care and Medical community.
- Responsible for resource planning and management (FTEs and phase IV budget) within Solid Tumors Therapeutic Area.
- May act as deputy to Medical Head Oncology

### **Essential Requirements:**

- Medical Degree is required
- from 5 to 10 years of experience in Pharmaceutical Industry in roles of increasing responsibility, with significant senior leadership experience in Medical Affairs
- Experience in global or regional roles is desirable
- Demonstrated experience and ability to lead, engage and inspire science-driven teams and organizations of more than 10 people
- · Strong cross functional collaborations skills
- Strong analytical and project management skills
- Experienced leading in a matrix environment
- Drive for innovation and continuous improvement
- Excellent communication skills
- Experience across multiple disease areas

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Company / Legal Entity IT08 (FCRS = IT008) Novartis Farma S.p.A.

Functional Area Research & Development

Job Type Full time

Employment Type Regolare

Shift Work No

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