

Regulatory Affairs Manager

Job ID REQ-10006524 Mai 10, 2024 Ukraine

Summary

Lead GMP Certificates (GMPC) recognition process: monitors validity of local GMP certificates recognitions, initiates prolongation and new GMPC recognitions, amendments of the product list, develop local dossier application forms, coordinate process with regulatory colleagues and other functions, inquire respective dossier from the production Site, coordinate cooperation with 3-d party regarding appropriate task description, order and performance; draft, agree and manage to receive necessary Power of attorney, coordinates respective expertise fee payments, submission and control of tasks' accomplishments. Directs the development of submission of product registration, progress reports, supplements, amendments, and/or periodic experience reports. Provides strategic product direction to teams on interaction and negotiates evidence with regulatory agencies. Interacts and negotiates with regulatory agency personnel in order to expedite approval of pending registration and answers any questions. Serves as a regulatory liaison on the project team throughout the product lifecycle. Ensures rapid and timely approval on of new drugs, biologics/biotechnology and/or medical devices and continued approved status of marketed drugs or medical devices. Serves as regulatory representative to marketing or research project teams and government regulatory agencies. Provides advice to development and/or marketing teams on manufacturing changes, line extensions, technical labeling, appropriate regulations and interpretations. Coordinates, reviews, and may prepare reports for submission.

About the Role

Major accountabilities:

- Is responsible for implementing regulatory strategy and managing operational activities for assigned medium regions.
- Provides input into global regulatory strategy and contributes to Regulatory Functional Plan (RFP) and Seed Document, or their equivalents, including identification of gaps or risks in global strategic plan for assigned regions.
- Partners with regions to align on regulatory strategy in order to fulfil business objectives -Implements RFP across assigned regions.
- Determines requirements and sets objectives for Health Authority (HA) interactions with DRA GPT representative and/or GTAL.
- Facilitates preparation and finalization of briefing books and contributes to preparation of summary documents.
- Develops and implements plans for timely response to HA requests and coordinates responses.
- May serve as local HA liaison depending on location (e.g., FDA or EMA).
- Drives coordination, planning, and submission of dossiers in assigned regions worldwide.
- Review of global dossier summary documents.

- Develops and implements plans to avoid/minimize clock stops during submission review.
- Reviews, approves and submits Clinical Trial Applications (CTAs) and Investigational New Drugs (INDs).
- Reviews and submits Risk Management Plans.
- May lead negotiations for regional approvals independently or with DRA GPT representative and/or GTAL.
- Responsible for facilitating timely submission and approval of dossier with HAs under the guidance of the DRA GPT representative and/or GTAL.
- Erroneous decisions result in critical delays and modifications to projects or operations; cause substantial expenditure of additional time, human resources, and funds; and jeopardize future business activity Contributes to and often leads the development of departmental goals and objectives.
- Reporting of technical complaints / adverse events / special case scenarios related to Novartis products within 24 hours of receipt -Distribution of marketing samples (where applicable)

Key performance indicators:

- Successful implementation of global regulatory strategy for timely submissions and approvals with the best possible labels based on available data.
- Identification of main HA issues -Participation in relevant regulatory Boards leading to valuable input from these Boards.
- Successful Participation in HA interactions to achieve business objectives.
- Adherence to Novartis policy and guidelines -Project & stakeholder feedback

Minimum Requirements:

Work Experience:

- Functional Breadth.
- Cross Cultural Experience.
- Operations Management and Execution.
- Project Management.

Skills:

- Regulatory affairs experience
- Detail Oriented.
- Drug Development knowledge
- Lifesciences.
- Negotiation Skills.
- Regulatory Compliance.

Languages:

• English.

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: https://www.novartis.com/careers/benefits-rewards

TEst Hello

Division

Global Drug Development

Business Unit

Innovative Medicines

Standort

Ukraine

State

Kyivska oblast

Site

Kyiv

Company / Legal Entity

UAPO (FCRS = CH024) NOPH SERVICES UKRAINE

Functional Area

Research & Development

Job Type

Full time

Employment Type

Regular

Shift Work

No

Apply to Job

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?

3/4

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: https://www.novartis.com/careers/benefits-rewards

The Novartis Group of Companies are Equal Opportunity Employers and take pride in maintaining a diverse environment. We do not discriminate in recruitment, hiring, training, promotion or other employment practices for reasons of race, color, religion, gender, national origin, age, sexual orientation, gender identity or expression, marital or veteran status, disability, or any other legally protected status. We are committed to building diverse teams, representative of the patients and communities we serve, and we strive to create an inclusive workplace that cultivates bold innovation through collaboration and empowers our people to unleash their full potential.

Job ID REQ-10006524

Regulatory Affairs Manager

Apply to Job

Source URL: https://dev1.novartis.de/de-de/careers/career-search/job/details/req-10006524-regulatory-affairs-manager

List of links present in page

- 1. https://www.novartis.com/about/strategy/people-and-culture
- 2. https://talentnetwork.novartis.com/network
- 3. https://www.novartis.com/careers/benefits-rewards
- 4. https://novartis4.wd3.myworkdayjobs-impl.com/en-US/Novartis_Careers/job/Kyiv/Regulatory-Affairs-Manager_REQ-10006524
- 5. https://www.novartis.com/about/strategy/people-and-culture
- 6. https://talentnetwork.novartis.com/network
- 7. https://www.novartis.com/careers/benefits-rewards
- 8. https://novartis4.wd3.myworkdayjobs-impl.com/en-US/Novartis_Careers/job/Kyiv/Regulatory-Affairs-Manager_REQ-10006524