Manager, Global Program Regulatory Manager (GPRM) Japan

Job ID REQ-10027926 Oct 31, 2024 Japan

About the Role

Major accountabilities:

- Assist developing innovative and high quality regulatory strategies to facilitate regulatory processes in development and ensure registration with optimized labels that contribute to health and welfare of the Japanese nation.
- Contribute to the regulatory activities in day-to-day operations for assigned TA area.
- Lead cross functional communication for preparing and finalizing Japanese labeling for new drugs.
- Lead regulatory related actions to maintain post marketing products in Japan.
- Establish strong relationship with the Japanese HA and obtaining high credibility in responsible projects.
- Ensure adherence to regulations, guidelines and global/internal procedures as required.
- Represent RA within specific internal discussions across line functions and external industry meetings.
- Mentor RA associates on drug development.
- Ensure adequate reporting of adverse events / technical complaint / compliance issue in accordance with company procedures.
- 100% timely delivery of all training requirements including compliance.

Key performance indicators:

- Achieve planned submission and approval in time in responsible projects.
- Obtain preferable outcomes of PMDA consultation in development phase projects for which the GPRM-J is responsible.
- No critical problem for maintaining post marketing product in responsible projects.
- Fulfil regulatory responsibilities in Japan to the GPT/GBT and RA subteam, and achieve registration with the best possible labeling.

Minimum Requirements:

Work Experience:

- Train and mentor RA members concerning drug development.
- Understand varied knowledge of Japan regulation.
- Possess extensive knowledge of MHLW/PMDA management, structures and organizations, and maintain trustful working relationship with MHLW/PMDA.
- Contribute to discussions on licensing conditions and integrate legal considerations into regulatory strategy.
- Possess extensive scientific knowledge of assigned TA/disease area, and facilitate scientific interactions

between experts relevant for drug development/maintenance.

- Address scientific issues across line functions and implement action plans.
- Define internal procedures for complying with effective regulatory requirements and enhancing quality and efficiency of the processes.
- Effectively negotiate with cross functional teams and lead an agreement in the optimal solution, and manage internal/external negotiation on development strategies and business critical issues.
- Excellent in effectively making presentation to clarify discussion items and raise key points to focus on in English.
- Contribute drug development planning by integrating expertise in the regulatory, legal and business environments.
- Possess extensive knowledge of global regulatory environment, and take appropriate actions to resolve issues identified in the projects that may negatively affect development strategy and progress.

Education:

- Degree in pharmacy, medicines, science, agriculture and/or pharmaceutical engineering discipline required. Advanced degree (Master Degree, PhD, etc.) preferred.
- Pharmacist license preferred.

Languages:

• Fluent English as business language.

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

Accessibility and Accommodation:

Novartis is committed to working with and providing reasonable accommodation to individuals with disabilities. If, because of a medical condition or disability, you need a reasonable accommodation for any part of the recruitment process, or in order to perform the essential functions of a position, please send an e-mail to diversityandincl.china@novartis.com and let us know the nature of your request and your contact information. Please include the job requisition number in your message.

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. You can follow us via Novartis Recruitment WeChat Official Account and Novartis Recruitment WeChat Video Account.

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-col/ture

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

Division

Development

Business Unit

Innovative Medicines

Location

Japan

Site

Head Office (Japan) (Pharmaceuticals)

Company / Legal Entity

JP05 (FCRS = JP005) Novartis Pharma K.K.

Functional Area

Research & Development

Job Type

Full time

Employment Type

Regular

Shift Work

No

Apply to Job

Job ID

REQ-10027926

Manager, Global Program Regulatory Manager (GPRM) Japan

Apply to Job

Source URL: https://jobapi.novartis.com/req-10027926-manager-global-program-regulatory-manager-gprm-japan

List of links present in page

- 1. https://jobapi.novartis.com/req-10027926-manager-global-program-regulatory-manager-gprm-japan
- 2. https://www.novartis.com/about/strategy/people-and-culture
- 3. https://www.novartis.com/sites/novartis_com/files/novartis-life-handbook.pdf
- 4. mailto:diversityandincl.china@novartis.com
- 5. https://talentnetwork.novartis.com/network
- 6. https://www.novartis.com/about/strategy/people-and-culture
- 7. https://talentnetwork.novartis.com/network
- 8. https://www.novartis.com/careers/benefits-rewards
- 9. https://novartis.wd3.myworkdayjobs.com/en-US/Novartis_Careers/job/Head-Office-Japan-Pharmaceuticals/Manager--Global-Program-Regulatory-Manager--GPRM--Japan_REQ-10027926-2
- 10. https://novartis.wd3.myworkdayjobs.com/en-US/Novartis_Careers/job/Head-Office-Japan-Pharmaceuticals/Manager--Global-Program-Regulatory-Manager--GPRM--Japan_REQ-10027926-2

3/4