



PR/095792 | Regional Regulatory Affairs Specialist

募集職種

人材紹介会社

ジェイエイシーリクルートメント シンガポール

求人ID

1575392

業種

福祉・介護

雇用形態

正社員

勤務地

シンガポール

給与

経験考慮の上、応相談

更新日

2026年04月28日 11:01

応募必要条件

職務経験

1年以上

キャリアレベル

中途経験者レベル

英語レベル

流暢

日本語レベル

無し

最終学歴

短大卒：準学士号

現在のビザ

日本での就労許可は必要ありません

募集要項

Job Title: Regional RA Specialist

Location: Singapore

About Us:

We are partnered with a leading pharmaceutical company committed to innovative healthcare solutions across Asia. They are dedicated to enhancing patient well-being through rigorous regulatory compliance and strategic pharmaceutical development.

They are seeking an experienced Regulatory Specialist based in Singapore to lead and support regulatory activities across the Asia & Arab region (excluding China & Japan). This role provides strategic regulatory advice, manages complex regional submissions and lifecycle activities, supports interactions with health authorities, and strengthens regulatory capability across local affiliates and cross-functional teams.

Key Responsibilities:

- Develop and execute regional regulatory strategies for new product registrations, renewals and major post-approval changes; anticipate risks and recommend mitigation to optimize approval timelines.
 - Lead coordination with local affiliates and Responsible Persons for submissions, product registrations, renewals and lifecycle management; review and validate ASEAN CTD / eCTD documentation.
 - Support interactions with Health Authorities by preparing scientific justifications, briefing materials and high-quality responses to queries.
 - Perform regulatory risk assessments for submissions and variations; track risk and propose actionable mitigation plans.
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- Monitor regulatory intelligence across APAC/ASEAN and the Arab region; analyze impacts and share insights to inform business decisions.

Qualifications:

- 2–5 years' regulatory affairs experience in the pharmaceutical industry, with regional exposure in APAC preferred.
- Strong understanding of APAC/ASEAN/ICH regulatory frameworks and country-specific requirements.
- Proven ability to manage complex regulatory tasks independently and provide strategic regulatory input.
- Experience preparing or reviewing ASEAN CTD / eCTD modules and supporting Health Authority interactions.

Interested applicants, please click **APPLY NOW**.

Please note, only shortlisted candidates will be contacted.

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会社説明