



Regulatory Affairs Assoc. Dir

外資系動物薬メーカーでの募集です。薬事申請のご経験のある方は歓迎です。

募集職種

人材紹介会社

株式会社ジェイ エイ シー リクルートメント

採用企業名

外資系動物薬メーカー

求人ID

1587583

業種

医薬品

会社の種類

外資系企業

雇用形態

正社員

勤務地

東京都 23区

給与

750万円 ~ 1600万円

勤務時間

09:00 ~ 17:30

休日・休暇

詳細は求人ご紹介時にご案内いたします。

更新日

2026年06月26日 20:00

応募必要条件

キャリアレベル

中途経験者レベル

英語レベル

ビジネス会話レベル

日本語レベル

ネイティブ

最終学歴

大学卒：学士号

現在のビザ

日本での就労許可が必要です

募集要項

【求人No NJB2327311】
【POSITION OVERVIEW】

The Associate Director of the Regulatory Affairs (RA) is responsible for the overall management of regulatory affairs activities including pharmacovigilance work at RA department at PDRA division Animal Health in Japan. The position requires a combination of scientific knowledge understanding of regulatory requirements strategic thinking and team leadership to ensure timely development of new products and regulatory affairs activities. The individual will ensure that the RA team develops and successfully executes project plans for timely product registrations.

The individual will motivate and encourage the RA Team to perform successfully and will mentor and coach team members to oversee their personal development in the organization.

The individual will closely communicate with members at the Product Development department to optimize the regulatory submission and get products approved.

【Reporting lines】

The RA Associate Director reports directly to the PDRA director

The RA Associate Director manages a team of 5-7 colleagues of RA department.

【PRIMARY ACTIVITIES】

Support PDRA director to optimize/maximize PDRA's activities by following ways:

Build PDRA as one team to have a seamless communication/collaboration

Review/advice Gap analysis done at PD department and create PDRA's one

Set up agreed product profiles among PD and RA.

Harmonize local requirements into the global development plan based on the well prepared development plan.

Liaise with Sales BU in the identification of market potential/requirements for new vaccines vaccine improvements and pharmaceutical products

Oversee the scientific quality of work within the unit to maintain and enhance reputation for excellence

Lead the RA team to ensure timely development of biological and pharmaceutical veterinary products in Japan in compliance with the general pharmaceutical guidelines and with local laws.

Ensure that the RA team develops optimized development plan by leveraging global data.

Oversee the collaborations with Contract Research Organizations (CROs). Ensure contract management and quality oversight of CROs.

Ensure compliance of organizational structures internal processes operational activities reporting including national and overseas pharmacovigilance requirements in line with the respective pharmaceutical law guidelines

Provide advice on research issues problems and trends having significant impact on RA activities of short and long term goals

Optimize the performance of RA colleagues by coaching motivating training and evaluating them properly in line with global and local development program including setting and reviewing individual yearly objectives and development plans

スキル・資格

【BACKGROUND REQUIREMENTS】

Following leadership behaviors are expected

- ・ Be Purposeful ・ Capturing the meaning of our work following our mission to improve people's lives and adhere to ethical behaviors.
- ・ Be empowering ・ Unfolding potential in self and others by providing guidance driving personal development and sharing enthusiasm.
- ・ Be collaborative ・ Working together across departments encouraging teamwork that respects diverse backgrounds and culture differences.
- ・ Be Results driven ・ Setting ourselves stretch performance target as well as establishing as establishing a feedback culture and taking responsibility.
- ・ Be innovative ・ Constantly experimenting and driving change initiatives. This includes being open to new ideas. Making quick decisions and taking risks.
- ・ Be future oriented ・ Continuously seeking best practice solutions and embracing opportunities of new technologies to shape the future of our business.

○Education

Veterinarian or PhD in Biological or Animal Sciences preferable.

○Experience

- ・ Minimum 7 years of relevant experience in a regulated industry preferably healthcare most preferably in the field of veterinary vaccines or pharmaceuticals.
- ・ Knowledge of Japanese Regulatory System preferable
- ・ Experience leading teams and developing/coaching people to maximize people/team capability.
- ・ Working in a matrix environment with multiple stakeholders at local and global levels.
- ・ Experience in planning and drafting budgets.

○Skills Knowledge and Competencies

- ・ Language ・ Japanese (fluent) English (professional level ・ verbal and writing)
- ・ Highly developed skills in negotiating influencing leadership and staff motivation.
- ・ Excellent communication and organizational skills.
- ・ Ability to work well with other departments in a multicultural team setting and matrix organization
- ・ Familiarity with animal diseases and animal husbandry techniques and ability to understand our product users including farmers and veterinarians.
- ・ Knowledge of Microsoft Office

会社説明

ご紹介時にご案内いたします