

グローバル企業・<mark>外資×ハイクラス転職</mark> 「語学力」を活かす転職なら、JAC Recruitment

[800~1500万円] Device Quality Assurance Manager/Sr. Manager

医療機器×医薬品 バイオベンチャーでの募集です。 メディカルGQP・GMP・品...

Job Information

Recruiter

JAC Recruitment Co., Ltd.

Hiring Company

医療機器×医薬品 バイオベンチャー

Job ID

1563536

Industry

Pharmaceutical

Company Type

International Company

Job Type

Permanent Full-time

Location

Tokyo - 23 Wards

Salary

8 million yen ~ 15 million yen

Holidays

【有給休暇】初年度 10日 入社1か月目から付与 【休日】完全週休二日制 土 日 祝日 夏季休暇 年末年始 ※詳細はオファー時...

Refreshed

November 27th, 2025 16:00

General Requirements

Career Level

Mid Career

Minimum English Level

Fluent

Minimum Japanese Level

Native

Minimum Education Level

High-School

Visa Status

Permission to work in Japan required

Job Description

【求人No NJB2337444】

Position Summary

This position will directly report to Head of Device Quality Assurance (QA) department and have a responsibility for handling and leading operation for quality management processes for the medical devices. It will provide opportunities to work closely with a multidisciplinary team in the group while mainly focusing on post market quality assurance matters management of licensees and product release for clinical trials.

Key Roles and Responsibilities

Lead post market activities including monitoring complaint trends maintaining risk management file performing products

and parts inspection among others.

- · Analyse and interpret data applying statistical methods to identify trends anomalies and opportunities for improvement.
- · Write review and maintain documentation of processes required for device quality management system.
- · Evaluate and manage suppliers both in Japan and overseas including collaborating with them to address quality challenges.
- · Work closely with a global multidisciplinary team to oversee feedback and complaint handling nonconformance control corrective and preventive actions change control supplier assessment among others.
- Maintain working knowledge of current regulations standards and guidance related to quality management systems and medical devices.

Required Skills

Desired Education Skills and Experience

- · A minimum of a bachelor's degree in science engineering or a related field.
- · At least 3+ years of experience and knowledge of relevant medical device industry. Candidates with more experience will be considered for a senior position.
- · Understanding of PMD Act relevant regulations and guidelines in Japan.
- · Thorough knowledge is expected of ISO 13485.
- · Knowledge of FDA 21 CFR Part 820 is a plus.
- · Experience in quality assurance operations listed below:
 - LQuality management system
 - LCAPA nonconformance feedback and complaint handling
 - └Product release and recall
 - LExternal and internal audits and
 - Linspection by regulatory authorities notified bodies and/or customers.
- · Comprehensive knowledge and practical experience in medical device risk management (e.g. ISO 14971) .
- · Hands on experience applying statistical tools and techniques to calculate sample sizes analyse data identify trends and support process improvements.
- Excellent verbal and written communication skills and ability to read write and speak Japanese.
- · Experience in communicating in English with business partners and ability to read write and speak in English is strongly preferred.

Company Description

ご紹介時にご案内いたします