



## 【800～1300万円】CMCマネージャー

外資製薬メーカーでの募集です。CMC薬事のご経験のある方は歓迎です。

### Job Information

**Recruiter**

JAC Recruitment Co., Ltd.

**Hiring Company**

外資製薬メーカー

**Job ID**

1591908

**Industry**

Pharmaceutical

**Company Type**

International Company

**Job Type**

Permanent Full-time

**Location**

Tokyo - 23 Wards

**Salary**

8 million yen ~ 13 million yen

**Work Hours**

08:30 ~ 17:15

**Holidays**

詳細は求人ご紹介時にご案内いたします。

**Refreshed**

May 28th, 2026 17:00

### General Requirements

**Career Level**

Mid Career

**Minimum English Level**

Business Level

**Minimum Japanese Level**

Native

**Minimum Education Level**

Bachelor's Degree

**Visa Status**

Permission to work in Japan required

### Job Description

**【求人No NJB2364696】**

- Responsible for CMC related regulatory affairs activities to obtain regulatory approval for biosimilars in Japan.
- Activities such as the preparation publication of REG CMC documentation for submissions to Health Authorities (HA).
- Interaction with HA on REG CMC questions to make new products and/or post marketed products successful.

( Major accountabilities )

- Lead CMC regulatory strategy with a focus on maximizing the business benefit balanced with regulatory compliance

- Lead submission activities ( planning authoring reviewing coordination submission ) for assigned projects/products.
  - Experience and capabilities are required to personally perform new submission and post marketed products.
  - Lead the submission by identifying required documents/ potential risks which could affect the defied timeline by assessing content and quality etc and by completing the application on time in line with a Global team • aligned project plan to obtain regulatory approval.
  - Authoring and reviewing high quality RA CMC documentation for HA submission ensuring alignment with applying agreed CMC global regulatory strategies and current regulatory trends and guidelines.
  - Prepare and communicate CMC Risk Management Assessments contingency plans and lessons learned on major submissions and escalate with management as appropriate.
  - Initiate and lead HA interactions and negotiations as appropriate; setting objectives preparing/reviewing briefing books coordinating and planning rehearsals and risk mitigation plans.
- 

## Required Skills

### ( Work Experience )

- Operations Management and Execution
- Cross functional collaboration

### ( Skills )

- Preparation of RA CMC related documents for biosimilar marketing authorization applications
- Change Control
- Cross Functional Teams
- Documentation Management
- Negotiation Skills
- Project Management
- Regulatory Compliance
- Risk Assessment
- Risk Management

### ( Languages )

- English and Japanese
- 

## Company Description

ご紹介時にご案内いたします