



## Senior/Advanced Project QA Professional

ノボ ノルディスク ファーマ株式会社での募集です。 メディカルGQP・GMP・...

### Job Information

#### Recruiter

JAC Recruitment Co., Ltd.

#### Hiring Company

ノボ ノルディスク ファーマ株式会社

#### Job ID

1578853

#### Industry

Pharmaceutical

#### Company Type

International Company

#### Job Type

Permanent Full-time

#### Location

Fukushima Prefecture

#### Salary

7 million yen ~ 15 million yen

#### Work Hours

08:50 ~ 17:10

#### Holidays

【有給休暇】有給休暇は入社時から付与されます 初年度 12日 1ヶ月目に付与（最高20日付与）【休日】完全週休二日制 年末...

#### Refreshed

May 17th, 2026 02:00

### 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 NJB2361383】

As a QA Professional you will play a key role in ensuring compliance with GxP regulations and maintaining the highest quality standards. Your responsibilities will include:

- ・ Providing QA oversight and support for projects in Koriyama factory in Japan to ensure timely delivery with the right quality.

- Participating in project and validation activities to resolve quality related issues.
  - Performing quality spot checks and approving validation documentation to ensure the highest quality level.
  - Reviewing and approving complex change control requests SOPs trend reports and validation related documents.
  - Preparing annual product reviews and facilitating validation and other complex internal trainings.
  - Acting as a coach during audits and inspections and ensuring compliance with regulations and corporate/local SOPs.
  - Providing proactive guidance to departments on quality related activities and ensuring that current Novo Nordisk procedures and health authority requirements are reflected in quality decisions.
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## Required Skills

### 【Your skills qualifications】

We are looking for a dedicated and experienced QA professional who can bring the following qualifications to the role:

- Minimum 5 years' QA experience or minimum 7 years' experience in QC/production or related roles ( healthcare/pharmaceutical industry preferred ) with a strong quality mindset and solid understanding of QA responsibilities.
  - Expert knowledge of quality management systems and cGMP production processes with deep understanding of validation in pharmaceutical manufacturing preferred.
  - Ability to make autonomous high - level decisions on complex quality issues and to perform effectively with minimal supervision.
  - Strong cross - functional leadership skills to drive process and team activities and to support others' ideas and actions.
  - Native level Japanese and business level English communication skills both written and oral.
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## Company Description

医療用医薬品、医療機器の開発、輸入・製造、販売