

# 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

# Pharmacovigilance Reporting Associate

ICONクリニカルリサーチ合同会社での募集です。 安全性情報 (臨床開発・製販後...

## 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

ICONクリニカルリサーチ合同会社

## 求人ID

1565696

## 業種

CRO

### 会社の種類

外資系企業

## 雇用形態

正社員

### 勤務地

大阪府

# 給与

450万円~600万円

# 勤務時間

 $09:00 \sim 17:30$ 

# 休日・休暇

【有給休暇】有給休暇は入社時から付与されます 入社7ヶ月目には最低10日以上 【休日】完全週休二日制 【休日】:土曜、日曜、祝...

# 更新日

2025年11月13日 15:01

# 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

流暢

## 日本語レベル

ネイティブ

# 最終学歴

専門学校卒

# 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 NJB2204577】

## Overview

- · Serve as safety reporting processor or lead for multiple safety reporting providing management support as designated.
- · Recognize exemplify and adhere to ICON's values which center around our commitment to People Clients and Performance.
- · As a member of staff the employee is expected to embrace and contribute to our culture of process improvement with a

focus on streamlining our

processes adding value to our business and meeting client needs.

- · Complete all departmental project activities accurately in accordance with ICON SOPs Study Specific Procedures regulatory requirements and client processes.
- Responsible for safety reporting or safety reporting intelligence activities on assigned projects working in a customer focused approach and an audit and inspection ready mindset.
- · Demonstrate skills pertaining to client management safety reporting project scope submission compliance quality and budget.

#### Detail

- · The following safety information case processing tasks related to clinical trials/post marketing of pharmaceutical products
- · Receipt of information on Adverse event triage numbering confirmation of details entry into database/QC
- · Creation of explanatory text for case course (Japanese and English) /QC
- · Primary evaluation of the necessity of reporting to the PMDA / QC of the evaluation details
- · Preparation of reports to PMDA/QC
- · Escalation coordination etc. to customers
- · Operations incidental to the above

\*Our Safety Reporting team will allow you to experience the ICCC study start up not just safety reporting. At first senior members will support you. You could expand your experience.

# スキル・資格

- · Experience required for any of the following
- · PV experience especially PMDA submission experience required.

Experience with ICCC is better.

- · 2+ years of CRA experience
- · Fluency in Japanese business level English

# 会社説明

1. 医薬品、医療機器、再生医療等製品、ワクチン等にかかる臨床開発、 市販直後調査、製造販売後調査、臨床研究等の受 託事業2. 労働者派遣事業