



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

1528012

業種

CRO

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

450万円 ~ 600万円

勤務時間

09:00 ~ 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03月19日 15:04

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