



【1000～1300万円】 Medical Monitor ※履歴書・英語CV必須<大阪・鹿児島>

株式会社新日本科学PPDでの募集です。医師のご経験のある方は歓迎です。

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

株式会社新日本科学PPD

求人ID

1584753

業種

CRO

会社の種類

外資系企業

雇用形態

正社員

勤務地

鹿児島県

給与

1000万円～1300万円

勤務時間

09:00～18:00

休日・休暇

【有給休暇】初年度10日4か月目から【休日】完全週休二日制 夏季休暇 年末年始 ■休日：年間122日（土日祝日、年末年始...）

更新日

2026年04月02日 15:07

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

大学卒：学士号

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 NJB2355107】

■What to do :

■Responsibilities: Project Related Medical/Safety Support:

◇ Minimize potential risk to PPD and clients by managing medical aspects of contracted tasks. This includes but is not limited to medical monitoring of all safety variables (AE laboratory abnormalities changes in patient medical status as well as inclusion/exclusion criteria evaluation prescribed concomitant medication for protocol restrictions and unblinding requests) .

Scope of work also includes discussion internally with medical monitors in Japan APAC and global and project team colleagues internally as well as with principal investigators and clients of all medical issues during the course of a study by proper medical judgment interpretation and decision.

◇ Medical review of serious adverse events: Ensure tasks delegated to medical monitors are properly executed. Adhere to applicable regulations and ICH guidance regarding clinical trials regulatory documents and safety issues. Adhere to client SOPs/directives and project specific WPDs for assigned projects. Adhere to PPD's corporate policies and SOPs/WPDs.

◇ Present PPD standard medical processes to clients at business development meetings investigator meetings and communicate with various medical communities to explore and expand PPD business.

◇ Provide medical consultation to team members and help manage protocol related medical questions. Communicate clearly with project team members and clients maintaining open communication to ensure all procedures are followed appropriately. Provide therapeutic training and protocol training on assigned studies as requested.

◇ Perform listing reviews as specified in the client contract and data validation manual including review of coding listings and/or full safety listings as well as use of Patient Profiles and other tools to assess for potential safety signal. This position will be assigned to APAC components of global or regional studies. Including but not limited to global/regional studies with Japan component.

◇ This person will also take care of a major responsibility of ICCG review therapeutic area training to clinical team in PPD SNBL supporting business development in PPD SNBL and clients requests including F2F visits to investigators KOLs and sites in Japan.

スキル・資格

■Required Qualifications

MD (Doctor of Medicine)

A certified license of a medical doctor

At least 2 years of clinical experience in a hospital setting (Experience in Neurology Oncology or General/Internal Medicine is a plus)

Ability to work effectively in both Japan based and regional/global environments

Fluency in both Japanese and English

■Prefferd Qualifications

Experience working with regulatory authorities and/or within the pharmaceutical industry

会社説明

臨床試験受託事業 (Contract Research Organization) ●臨床第1 4相試験におけるモニタリング●国内、アジア、およびグローバルの臨床試験のプロジェクトマネジメント●生物統計解析、データマネジメント●ファーマコビジランス (安全性監視業務) ●PMDA対面助言に関わる支援を含む薬事業務●メディカルライティング●GCP QA業務 (Investigator Site Audit, Vendor Audit、等)