



## <研究開発・メディカルアフェアーズ統括本部>Project Statistician/ Principal Statistic...

日本イーライリリー株式会社での募集です。統計解析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

日本イーライリリー株式会社

#### 求人ID

1570330

#### 業種

医薬品

#### 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

600万円～経験考慮の上、応相談

#### 勤務時間

08:45～17:30

#### 休日・休暇

【有給休暇】初年度 10日 2か月目から 【休日】完全週休二日制 年末年始 完全週休2日制（土・日曜日）、祝日、クリスマス、年...

#### 更新日

2026年01月22日 16:00

### 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

ビジネス会話レベル

#### 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 NJB2348617】

#### 職務内容

The Project statistician develops or assists in the development of protocol designs clinical plans and data analysis plans in collaboration with physicians veterinarians and/or medical colleagues. The Project Statistician is responsible for working with research associates and scientists to establish a reporting database and for analyzing data for these types of studies.

**[Clinical Project Statistician]**

Provide strong statistical leadership in the process of drug development.

**[Real World Analytics Project Statistician]**

Provide input and statistical leadership for post launch activities including Health technology assessment (HTA) real world evidence (RWE) research post market safety studies medical affairs publications to maximize the value of our products in collaboration with cross functional and cross regional partners.

**主な職責/Primary responsibilities****Statistical Trial Design and Analysis****[Clinical Project Statistician]**

- ・ Operate in collaboration with study personnel to provide input on study protocol design studies and write protocols for the conduct of each study.
- ・ Assist in or be accountable for selecting statistical methods for data analysis authoring the corresponding sections of the protocol and conducting the actual analysis once a reporting database is created.
- ・ Collaborate with data management in the planning and implementation of data quality assurance plans.
- ・ Maintain currency with respect to statistical methodology to maintain proficiency in applying new and varied methods and to be competent in justifying methods selected.
- ・ Participate in peer review work products from other statistical colleagues.

**[Real World Analytics Project Statistician]**

- ・ Lead statistical activities with the development of protocols analysis plans and analysis execution and interpretation for RWE research and post market safety studies.
- ・ Provide technical statistical leadership for network meta analysis/indirect comparisons economic modelling and dossier development for Health technology assessment (HTA) .
- ・ Ensures high quality statistical support for projects through the oversight of external suppliers and provides input into outsourcing strategy and processes.
- ・ Influences cross functional team members regarding appropriate research method.

**Communication of Results and Inferences**

- ・ Collaborate with team members to write reports and communicate results.
- ・ Assist with or be responsible for communicating study results via regulatory submissions manuscripts or oral presentations in group settings as well as for communicating one on one with key customers and presenting at scientific meetings.
- ・ Respond to regulatory queries and to interact with regulators.

**Therapeutic Area Knowledge**

- ・ Understand disease states in order to enhance the level of customer focus and collaboration and be seen as a strong scientific contributor.

**Regulatory Compliance**

- ・ Perform work in full compliance with assigned curriculum (s) and will be responsible for following applicable Corporate Medical local and departmental policies procedures processes and training

**スキル・資格****■Basic Requirements:**

- ・ M.S. or Ph.D. in statistics or biostatistics

**■Other Information/Additional Preferences:**

- ・ Rich experiences in healthcare field as a statistician with science background such as mathematics or epidemiology.
- ・ Statistical and methodological knowledge in clinical development epidemiology or related field.
- ・ Regulatory knowledge of clinical trial methodology and statistics.
- ・ Ability to build relationships with individuals and teams.
- ・ Good communication and presentation skills in both English and Japanese

**会社説明**

医療用医薬品の輸入・製造・販売