



【1000～1500万円】 【External Quality】 Associate Quality Director

アストラゼネカ株式会社での募集です。 メディカルGQP・GMP・品質保証・品質...

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

アストラゼネカ株式会社

求人ID

1522692

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

1000万円～1500万円

勤務時間

09:00～17:15

休日・休暇

【有給休暇】初年度10日1か月目から 【休日】完全週休二日制 年末年始 私傷病休暇、サパティカル休暇、ボランティア休暇、慶...

更新日

2025年02月20日 16:14

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 NJB2280811】

The incumbent is responsible for the Quality management of assigned External Suppliers within Procurement Quality AsiaPac teams. They are responsible for all Quality activities that directly support execution of Quality Management of Suppliers for assigned suppliers. This includes but is not limited to the Quality System oversight and/or performance of the following activities: change control product quality complaint S L complaint and deviation investigations quality issue management and escalation Quality Agreements (establishment and maintenance) between AZ and External Suppliers

and between EQ and AZ Operations Sites.

Within the External Quality (EQ) organization the job holder is responsible for the oversight and ownership of Quality System (s) . They will support the Quality Professionals involved in Quality Supplier and Product Supply Chain Management within the assigned categories.

In addition the preparation and submission of periodic Supplier Quality Assessments Regulatory Agency interactions and serving as Quality leaders on NPI new supplier introductions In Licensing strategic sourcing projects process optimization and product transfer projects as these relate to Quality Supplier management are within the scope of this role. Regulatory Agency interaction includes preparation for and management of Regulatory Agency inspections at External Suppliers and AZ sites (when External Suppliers are assessed) .

スキル・資格

【経験/Experience】

(歓迎/ Nice to have)

- Experience working in a PCO/PET organization or Lean/Six Sigma training.
- Multi site / multi functional experience
- Proven experience in Quality Assurance or combination of Quality and Technical
- Masters Degree in Quality Assurance/Regulatory Affairs or other advanced scientific field

【資格 /License】

(必須 /Mandatory)

Bachelor degree in a science / technical field such as Pharmacy Biology Chemistry or Engineering (Note: there may be specific additional requirements depending on the regulations in each country) . Proven broad experience in either the pharmaceutical operations environment or pharmaceutical Quality Assurance role.

【能力/ Skill set】

(必須/ Mandatory)

- Strong demonstrated knowledge of cGMPs Quality Systems and the pharmaceutical supply chain environment. Also strong understanding of industry standards such as Pharmacopeia ISO standards etc.
 - Excellent oral and written communication skills [English and local language (s)]
- pecifically required essentials for Career Level E:
- Demonstrated experience working cross functionally and managing significant improvement initiatives (e.g. project management skills)
 - Strong problem solving skills
 - Strong negotiating/influencing skills
 - Ability to work independently under his/her own initiative.

【その他/ Others】

(必須/ Mandatory)

- Ability to travel nationally and internationally as required approximately 10% of their time.

(歓迎/ Nice to have)

- PCO members or equivalent team
- Category/Sub Category Team members
- Quality and other support groups within or across sites
- Regulatory Affairs (including GQO CMC RC)
- GQO
- Global Supply Managers/Directors
- Pharmaceutical Technology and Development and Pharmaceutical Sciences

会社説明

医療用医薬品の開発、製造及び販売