

Title: Senior Manager/Associate Director, Quality Assurance

Job Summary:

Reporting to the Sr. Director, Quality Assurance, the Senior Manager/Associate Director, Quality Assurance is responsible for the execution and administration of GXP Quality Systems as they pertain to applicable cGMP, GLP, and GCP regulations. The Senior Manager/Associate Director will work with cross functional teams to maintain the internal quality systems associated with GxP activities as well as interfacing with CMO/CROs.

Responsibilities:

- Perform Quality & Compliance oversight for manufacturing, supply chain, clinical, & nonclinical activities
- Assist in build-out, management and continuous improvement of the QMS and SOP system
- Assure consistency in achieving product quality and compliance across multiple CMOs and CROs
- Assist in the execution of QA systems to support GxP from both an internal and external perspective
- Plan audit calendar, perform internal/external audits
- Provide "man-in-plant" observation/feedback at key manufacturing milestone activities
- Review and negotiate Quality Agreements; review MSA/SOW to ensure compliance with the QA agreement
- Develop and conduct training
- Write, review and approve Standard Operating Procedures (SOP) and Forms for the organization
- Review batch records and associated documentation, i.e., deviations, OOSs, investigations, specifications, etc.; lead internal investigations.
- Trend and report QA related information (deviations, investigations, CAPA) both internally and from external activities
- Assist with other GXP programs and reviews as needed

Qualifications:

- BA/BS in Life Sciences
- Minimum of 10 years' experience in GxP regulated industries
- Extensive knowledge and experience in GXP and QA principles, practices & industry standards
- Experience using risk-based principles & decision making to ensure compliance at all stages of development
- Experience auditing contract organizations, auditor certification preferable
- Experience working in small biopharmaceutical company and/or experience working within a virtual manufacturing company utilizing a number of contract manufacturing organizations is preferred
- Excellent attention to detail, project and time management skills, and the ability to manage multiple priorities with aggressive timelines
- Ability to travel up to 25% of the time
- Excellent written and verbal communication skills
- Ability to work effectively/congenially both independently and with a multi-disciplinary team

If you would like to be considered for a job on the Concert team, send your resume in the body text of an e-mail (no attachments please) to: careers@concertpharma.com.

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