

Als kompetenter Personaldienstleister bringt PERSONALHANSA seit 40 Jahren Menschen in Arbeit. Mit Engagement und Expertise gelingt es uns die Interessen der Bewerber und unserer Kunden bestmöglich in Einklang zu bringen. PERSONALHANSA ist Mitglied der GVP-Tarifgemeinschaft. Berufsanfänger und Berufserfahrene erhalten bei uns, neben sehr guten Verdienstmöglichkeiten, attraktive und sichere Orientierungsmöglichkeiten auf dem sich ständig ändernden Arbeitsmarkt.

Wir suchen für unseren Kunden in der Metropolregion Rhein-Neckar einen

## **GMP Compliance Auditor (m/w/d)**

### **Hauptaufgabengebiet:**

- ▶ Carryout job duties independently, including interpret, explain and apply applicable current regulations, guidelines, policies, and procedures for pharmaceutical products, medical devices, and regulated studies
- ▶ Independently plan and conduct internal system audits and external GxP supplier audits
- ▶ Maintain approved supplier list, global audit schedule and participate in supplier management processes
- ▶ Gathers internal and external audit metrics and presents to QA management for trend analysis
- ▶ Recommend plan of action for satisfactory resolution of quality and regulatory compliance issues
- ▶ Provide guidance and training on GxP regulations and guidelines to GxP auditors and functional areas personnel
- ▶ Assist GxP QA Management on the collaboration with the Inspection Management group on external audits by regulatory agencies and customers
- ▶ Develop / maintain and update departmental systems, procedures and records pertinent to position responsibilities
- ▶ Expected to elevate any issues to management, as necessary, in meeting these responsibilities. Leads goals with cross-functional or broader scope. Resolves project team issues with minimal oversight

## Qualifikationen:

- ▶ Bachelor's degree preferably in life sciences or engineering
- ▶ Demonstrated strong leadership competencies, proficient level of technical capabilities, and independence. Proven track record utilizing core & technical competencies
- ▶ Thorough understanding of international GMP regulatory standards
- ▶ 5+ years experience in function or related fields such as:
  - Quality Assurance / Regulatory Affairs
  - Pharmaceutical / Device / Healthcare Industry
- ▶ Quality Assurance auditing experience (preferred)
- ▶ Laboratory experience (preferred)
- ▶ An equivalent combination of education and experience may be accepted as a satisfactory substitute for the specific education and experience listed above
- ▶ Accreditation by a professional body is desirable (e.g. American Society for Quality (ASQ) Certified Quality Manager (CQM) and/or Certified Quality Auditor (CQA))

## Unsere Leistungen:

- ▶ Vergütung mindestens nach GVP-Tarifvertrag
- ▶ Einsatzbedingte Zulagen und/oder Branchenzuschläge
- ▶ Urlaubs- und Weihnachtsgeld
- ▶ Arbeitszeitkonto auf Wunsch erst ab der 41. Wochenarbeitsstunde
- ▶ Dauerhaft Einsätze – keine Tageseinsätze
- ▶ 25 % Zuschuss zum Deutschlandticket
- ▶ Übernahmeoption bei unseren Kunden
- ▶ Intensive Betreuung / Beratung bei der beruflichen Weiterentwicklung

Ihre Bewerbung senden Sie bitte an: [bewerbung@personalhansa.com](mailto:bewerbung@personalhansa.com)