



## **Clinical Project Manager (Contractor)**

**Location:** London, UK

**Contract Length:** 6 months

**About Us:** At Nightstar, our mission is to maintain and restore sight in patients with inherited retinal diseases. We are a clinical-stage company focused on developing and commercializing a pipeline of novel and potentially curative, one-time retinal gene therapies for patients suffering from rare inherited retinal diseases that would otherwise progress to blindness, and, for which, there are no currently approved treatments.

**Job Purpose:** We are seeking a Contract Clinical Project Manager for the Company's London office. This is an excellent opportunity for someone with strong organizational skills and the ability to work on a variety of tasks and projects simultaneously with minimal supervision. In addition, the successful candidate must have strong project management experience, good judgment, be flexible when different tasks arise, and be detailed oriented.

### **Job Responsibilities:**

- Working with the clinical team to support design, development, execution, and delivery of the clinical study in accordance with the clinical development plan/strategy and timelines
- Collaborate closely with the Program Lead, working on execution and oversight of sponsored clinical trials
- Leading in the oversight of CRO activities and other clinical vendors to ensure study quality meets Nightstar and regulatory requirements
- Supporting the Program Lead and medical writers to initiate protocols, study reports, Investigator Brochures, and other key clinical documents
- Responsible for oversight of identification and selection of investigator sites
- Managing escalation of study related issues and communicates as appropriate with management and other R&D functions
- Ability to anticipate potential study issues and to prepare contingency plans with minimal oversight
- Collaborating with the cross functional teams on clinical activities (drug supply, resolution of data queries, etc.) and on selection and management of clinical vendors
- Maintaining knowledge of therapeutic area, current medical practice and pharmaceutical regulations in order to ensure best practice across all activities

### **Critical Competencies:**

- A minimum of 8 years of industry experience in the biotechnology/biopharmaceutical industry
- A minimum of 3 years of Project Management experience, including experience running global studies
- Experience in clinical operations and development, preferably Phase III
- Working with the Program Lead and medical writers to initiate and write protocols, study reports, IBs, and other key clinical documents
- Working with CROs and overseeing vendors, preferably in Phase III studies
- Working with multi-national studies

### **Behavioral and Interpersonal:**



- Exceptional attention to detail and excellent organizational skills with a desire to roll up one's sleeves
- Excellent oral and written communication skills
- Ability to thrive in a dynamic and fast-paced environment
- Ability to prioritize duties and manage multiple matters from start to finish with minimal supervision with a demonstrated ability to lead change and make independent decisions
- Ability to effectively and positively work with executive-level management
- Ability to handle highly confidential and sensitive materials and information with complete discretion and having good judgment in working with clients, and occasionally under ambiguous or challenging circumstances
- A dynamic self-starter with a positive attitude and strong influencing skills

**Education Requirements:**

- Bachelor's degree, nursing background is also acceptable