

Senior Clinical Research Associate at BeiGene, New Zealand

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The CRA is responsible to deliver data within timelines and required quality standards, responsible for adherence to monitoring procedures in accordance with GCP and ICH, local regulations and SOPs



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BeiGene continues to grow at a rapid pace with challenging and exciting opportunities for experienced professionals. When considering candidates, we look for scientific and business professionals who are highly motivated, collaborative, and most importantly, share our passionate interest in fighting cancer.

Job Description:

Purpose of the Job:

- The CRA performs monitoring activities related to initiation, conduct (recruitment, quality data collection) and timely completion of Oncology and Pharma clinical trials within the country.
- The CRA is responsible to deliver data within timelines and required quality standard, responsible for adherence to monitoring procedures in accordance with GCP and ICH, local regulations and SOPs.

Main Responsibilities:

- Conducts monitoring (pre-study, initiation, routine monitoring and closeout visit), if required
- · Conducts co-monitoring visits, if required
- Conducts Quality Oversight Visits (QOV), as requested
- Completes monitoring visit/ QOV reports timely
- Assists with investigator/site identification
- · Assists site to prepare Ethics Committee submissions
- Facilitates clinical trial site contract and budget negotiation
- Manages site queries and communications
- · Assists in managing clinical trials, if required
- · Establishes regular lines of communication with sites and COMs
- Provides protocol and related study training to assigned sites
- Evaluates the quality and integrity of site practices escalating quality issues as appropriate
- Manages site performance by tracking regulatory submissions, recruitment, case report form (CRF) completion, and data query resolution

Qualification Required:

- Bachelor's level degree or above in life sciences, pharmacy, nursing or medical
- Understands clinical trial processes with a thorough knowledge of ICH and associated regulatory guidelines
- 4 years or more monitoring experience in the Pharmaceutical or CRO industry
- · Excellent communication and interpersonal skills
- · Excellent organizational skills and ability to prioritize and multi-task
- Fluent in English (writing and speaking)
- · Applicant must have full working rights in New Zealand to be considered

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