Intrinsik Corp., is a North American consulting firm focusing on the toxicology and regulatory challenges associated with the development of new products (pharmaceuticals, biologics, consumer products, natural health products and cosmetics). We are continuing to grow, and will always welcome candidates with strong scientific skills, creativity and enthusiasm to join our team.

Intrinsik has an opening for a **Scientific Editor** who is looking for an opportunity to work in an environment that truly values scientific excellence, creativity, and initiative. The Scientific Editor will review and revise nonclinical, clinical, and regulatory documentation according to client and company guidelines. The Scientific Editor provides accuracy, consistency, and contributes to cross-functional working groups as needed to facilitate the preparation of reports, and the production of various regulatory dossiers.

We aim to offer our employees an environment that encourages professionalism, creativity, independence, and continual learning. The assets of any knowledge-based company are its people, and we believe strongly in investing in those assets by offering training and mentoring of our staff. Like all modern successful organizations, we are committed to growing and advancing our employees' careers by providing them with new responsibilities and opportunities within the company.

**Responsibilities include:**

* Providing quality control and reviewing clinical study protocols and clinical study reports (Phase 2/3 studies), and other regulatory submission documents (e.g., CTD Modules 2.5 and 2.7, and Integrated Safety/Efficacy Summaries, Investigator’s Brochures, and briefing documents for regulatory meetings).
* Analyzing and interpreting scientific/clinical data.
* Quality control of systematic scientific literature searches, including tabulated and written summaries of publications.
* Overseeing quality control for nonclinical, clinical and regulatory projects.

**The successful applicant(s) would ideally have the following qualifications and attributes:**

* Degree in the life science.
* Demonstrated high degree of independence and proficiency with scientific writing and attention to detail.
* Excellent interpersonal, verbal, and written communication skills.
* Understanding of the drug development process and applicable industry guidelines and regulations (e.g., ICH).
* Minimum of 3 years of quality control experience in the pharmaceutical industry.
* Ability to multi-task (multiple projects/issues) and prioritize workload within a deadline-oriented, fast-paced environment.
* Ability to work both autonomously and collaboratively with a team.
* Extensive knowledge of English grammar with proven written and spoken English.

Interested candidates may submit a **cover letter** and resume viae-mail to Heather Wilson at hwilson@intrinsik.com. We thank all candidates for applying; however, only those considered for an interview will be contacted by Human Resources.