Job Details

Job Title Lawson, Clinical Research Associate - Lawson Clinical Research

Location Victoria Hospital

Regular/Temporary Regular

Job ID 60768

Full/Part Time Full-Time

Favorite Job

Posting Period

March 26, 2018 Open: Deadline: April 1, 2018

Non-Union

Different terms and conditions of employment may apply to externally funded positions

Department Name



Lawson Health Research Institute - LHR Clinical Trials Services

Lawson Health Research Institute (Lawson) is the research institute of London Health Sciences Centre and St. Joseph's Health Care London. As one of Canada's top ten research institutes, we are committed to furthering scientific knowledge to advance health care around the world

As a member of the Lawson Clinical Research Services team, the Research Associate will collaborate with Investigators and health care teams to assume responsibility for the overall patient management and coordination of several clinical studies

Studies may include all programs supported at the city wide sites for LHSC and SJHC. They will include industry-sponsored, cooperative group sponsored (such as CIHR) as well as Investigator initiated trials.

This position will provide an excellent opportunity for a dynamic individual with demonstrated organizational and communication skills.

Responsibilities include, but are not limited to, study start up, recruitment of study participants (e.g. identify and screen potential subjects, obtain informed consent); coordination of patient visits schedules as per study protocol; execution of all aspects of study visits (e.g. assessment adverse events, monitoring safety, medication, questionnaires, sample collection, (may including processing and shipment of samples according to clinical protocol), provides clinical care for patients participating in clinical trials and the implementation and coordination of all aspects of data collection and source documentation, as per LHSC policy and ICH/GCP guidelines. This position will require professional communication both written and verbal between all clinical trial stakeholders

Rate of Pay: To commensurate with experience

Hours of work: 37.5 hours per week

Qualifications

- Successful completion of Bachelor's Degree in Health Sciences or related field of study
- Minimum of two years clinical research experience with industry sponsored clinical trials an asset Designation in SOCRA, ACRP an asset
- Diploma or Certificate in Clinical Trials Management is an asset
- Depoints of extinated in Continual Trials with a Brasset Philebotomy skills and Transportation of Dangerous Goods/ International Air Transport (TDG/IATA) are an asset Demonstrated knowledge of current regulations and guidelines for conducting clinical trials (ICH GCP, Health Canada Food and Drug Regulations (Part C, Division 5), USA FDA 21 CFR, TCPS2 Tri-Council Policy Statement, and privacy legislation (PIPEDA, PHIPA)
- Knowledge of local REB requirements and Lawson approval process
- Demonstrated knowledge of research medication management and accountability
- Well-developed patient assessment and evaluation skills

 Demonstrated ability to communicate effectively and professionally with study participants and co-workers
- Demonstrated cooperation in a team environment
- Demonstrated ability to plan, prioritize, execute and manage several research studies

 Demonstrated computer proficiency in Microsoft Office and experience with electronic data entry (eCRFs) and databases
- Excellent verbal and written communication skills in English
- Highly motivated and self-directed
- Demonstrated knowledge of and commitment to the principles of patient and family centered care
- Demonstrated knowledge of and commitment to patient and staff safety at LHSC
- Demonstrated ability to attend work on a regular basis

London Health Sciences Centre foster a culture of patient and staff safety whereby all employees are guided by LHSC's Mission, Vision, Values and Code of

We are committed to providing a safe, healthy and inclusive work environment that inspires respect. LHSC encourages applications from persons with disabilities and we are committed to providing accommodations upon request.

As part of the assessment process applicants may be required to complete a written examination or test. Please be advised that an internal reference check may be conducted as part of the selection process.

Your interest in this opportunity is appreciated. Only those applicants selected for an interview will be contacted