Toxicological Pathologist

Company Overview

Accelera is a contract research lab that provides integrated services for discovery and preclinical drug development. Accelera offers a unique alternative by integrating nonclinical disciplines in pharmacology, drug metabolism and pharmacokinetics, toxicology, and pathology in order to helping our Partners expedite their preclinical drug development with exceptional safety assessment services, state-of-the-art facilities and expert regulatory guidance.

We are currently looking for a Senior/Principal Toxicological Pathologist to be involved in the drug development process with passion for science and quality.

Position

Accelera has a full-time opening for a Toxicological Pathologist with demonstrated expertise in drug development. The successful candidate will have a broad knowledge of toxicological pathology across multiple species to provide expert pathology support working in a collaborative environment on projects throughout the discovery and development continuum.

Qualified individuals will also participate in pathology oversight for regulatory toxicology studies, including the evaluation of clinical pathology data, primary histopathology evaluations for in-house studies, and pathology peer review for GLP studies.

Key Responsibilities

- Conduct gross and microscopic examination of animal tissues from toxicology and investigative studies
- Diagnose and interpret compound-related effects in animals including assessment of cause of death, target organs of toxicity and reversibility of toxicity by providing expert primary pathology reads on exploratory and definitive toxicity studies
- Provide concise written pathology reports that accurately and completely reflect the data collected and the impact on drug development, drug registration and human safety
- Conduct pathology peer reviews of GLP studies
- Summarize experimental results for presentations both within and outside of the Company

Qualifications

- Doctor of Veterinary Medicine (DVM) or equivalent
- Board certification in veterinary pathology (ECVP or ACVP) (or board eligible)
- At least 3-5 years of experience interpreting and reporting pathology findings for regulated (GLP) studies, preferably in a pharmaceutical or biotechnology company or a CRO conducting studies for pharmaceutical or biotechnology companies
- Strong, effective written and oral communication skills including the ability to organize and clearly present complex data and concepts
- Computer skills, commensurate with essential functions, including the ability to learn a validated system.
- Ability to work in a fast-paced environment, manage multiple projects, demonstrate consistent productivity, and meet project timelines

If your background and personal experience fit the profile, please send your complete application, including a brief description of your experience and your CV and specifying in the subject “TOXICOLOGICAL PATHOLOGIST” to the following address:

jobposting@nmsgroup.it
Please add to your CV the statement “In compliance with the EU Regulation n. 679/2016 (GDPR) and with the Italian Law n. 196/2003, I hereby authorize the recipient of this document to use and process my personal data for the purpose of recruiting and selecting staff.”

The Company is an equal opportunity employer and prohibits unlawful discrimination based on race, color, religion, gender, sexual orientation, gender identity/expression, national origin/ancestry, age, disability, and marital status.