

MICHAEL AKINS, PhD

Dr. Akins has over 20 years of experience in academic research where he applied his broad training in cell and molecular biology to the study of animal models of nervous system disorders. During that time, he worked in lead and collaborative roles to produce numerous manuscripts, book chapters, and presentations. He recently transitioned to regulatory writing and is gaining experience developing documentation for regulatory submissions.

ARCHANA PARASHAR, PhD

Dr. Parashar is a biomedical scientist with a diverse background and more than 10 years of experience in the areas of drug discovery, vaccine development, in vitro diagnostics, biomedical product manufacturing, and bioinformatics, as well as in scientific writing, including authoring research publications, grants, reports, and communications. Having worked as a scientific lead, cross-functional liaison, and project manager, Dr. Parashar is goal oriented and a honed team player. She is passionate about regulatory writing.

MRS. PAMELA ROGERS, MS

Mrs. Rogers is a pharmaceutical industry professional with experience in laboratory analysis, client management, project management, and laboratory inventory management. Her previous roles include performing high throughput screening in support of drug discovery, executing bioanalytical laboratory analysis (small and large molecule), as well as conducting academic research work in molecular biology. Mrs. Rogers' technical writing experience includes writing, editing, and review of laboratory protocols, sample analysis plans, methods, and study reports for bioanalytical studies, as well as completion of a graduate thesis. She also has prior experience with quality control of generated laboratory data.

JENNIFER SMITH, PhD

Dr. Smith has over 10 years of experience as a research scientist. She is experienced in drafting, editing, publishing, and reviewing manuscripts and has independently managed a variety of regulatory documents vital to basic science research. Her area of expertise is in systems and molecular neuroscience with an emphasis on neuromodulation. Dr. Smith is a new addition to the regulatory writing team and is passionate about supporting clinical innovation and promoting public health.

TRICIA WRIGHT, PhD

Dr. Wright has been involved in biomedical scientific research for over 12 years, with bench research experience and a background in molecular and cancer biology. She has experience in writing first-author peer-reviewed scientific manuscripts, writing peer reviewed review articles, and successfully writing funded scientific grants and fellowships. She has also written and developed abstracts, posters, and presentations.