

CASE STUDY

Medical Writing

Client: *Biopharma Startup Client*

SITUATION

A startup biopharmaceutical company asked Pearl for assistance with medical writing to prepare for needed publications in support of an upcoming IND approval and launch. The client required assistance composing several pharmacokinetics publications, abstracts and posters.

SOLUTION

Pearl provided a team of medical writers to help the company for 4 months producing a total of 8 publications and 8 abstracts/posters. Pearl led all medical writing activities and worked across a cross functional team at the client to project manage the project including efforts to secure needed references and raw data. Pearl also worked to ensure all publications met journal publishing requirements.

RESULT

Pearl worked closely with the client team, and provided high quality draft publications, abstracts and posters. As a result of Pearl's work, the client exceeded project deadlines, and was extremely pleased with the quality of the writing, and is currently seeking further work with Pearl.

Pearl Pathways is a comprehensive life science product development services company. Every day we strive to provide our customers top quality service, unyielding ethics, and efficient services through our team of experts. Pearl Pathways supports biopharmaceutical, medical device, and diagnostic companies as well as life science service providers with clinical, regulatory, and quality compliance needs. Our full-service central IRB supports all aspects of human research.

Clinical: Pearl Pathways' expertise in medical practice, science, ethics, and clinical research supports all phases of clinical development. Our expert consultants help implement customized solutions that drive efficient clinical research programs.

Regulatory (drug): Our consultants draw on over 20 years of scientific, industry, and FDA experience to translate your data into strategic drug development pathways, providing guidance from molecule to market while navigating a complex regulatory environment.

Regulatory (device): Delivering superior end-to-end product development and commercialization strategies, our consultants provide regulatory guidance that adds decades of industry and FDA regulatory expertise to your team, helping you balance risk, cost, and time to market.

Quality: From risk-based quality system development to gap analysis and remediation, our partnership helps reduce operational burdens, manufacturing impacts, regulatory agency warnings, and costly recalls—allowing you to focus on ongoing product quality assurance.

IRB: Our accredited independent review board ensures quality research, unyielding ethics, and human subject protection with individualized Expedited and Full Board reviews.

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