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Mark Slama Named New G.M. for Bio-Kinetic Clinical Applications

(September 16, 2008) QPS/Bio-Kinetic announces that Mark Slama will take over responsibilities as general manager of Bio-Kinetic Clinical Applications, the company's Phase I facility. Slama had formerly been director of clinical operations at Qualia Clinical Services where he more than doubled the size of his division and substantially increased revenue.

"We're proud to add someone of Mark's caliber to our team," Bio-Kinetic Clinical Applications CEO Dale Bourg said. "His knowledge of the CRO industry is both broad and deep, and his experience in the full range of operations has given him extraordinary insight.

"Prior to joining Qualia, Slama had spent over 17 years at MDS Pharma Services, starting as a quality control technician and moving up through the company with roles in study coordination, facility and staff scheduling, clinical research, and support services management. He left the company as manager of the global bids and contracts department, a post he had held for almost eight years.

"Mark's experience includes directly managing or monitoring over 200 Phase I trials as well as Phase II and III trials in the areas of oncology, pain management, and renal and allergic retinitis," Bourg said. "We're certain his depth of knowledge will prove to be of great value to our clients."

Slama holds a BA degree in organizational management from Concordia University and is a veteran of the Nebraska Army National Guard.

About Bio-Kinetic

Bio-Kinetic, www.qpsbiokinetic.com, is a drug development services company that designs and conducts Phase I clinical studies for branded pharmaceutical, bio-technology, and generic drug companies. The company owns and operates a 240-bed Phase I facility in Springfield, Missouri which positions Bio-Kinetic as one of the larger independent Phase I facilities in the country. The Company is known for its clinical quality, having developed an outstanding reputation for study designs, recruiting and screening study participants, and collecting and reporting clinical data.

About QPS

Founded in 1995, QPS, www.questpharm.com has bioanalysis, DMPK, and Translational Medicine facilities at its Newark, DE, headquarters and a laboratory in Taipei, Taiwan. Regional sales offices are maintained in California, Connecticut, Massachusetts, Ohio, Pennsylvania, and Texas. QPS is a GLP-compliant CRO that supports discovery, preclinical and clinical drug development. It has been providing quality services in bioanalysis (LC/MS/MS, LBA, Hybridization-LBA), preclinical and clinical DMPK (pharmacokinetics, metabolism, autoradiography), translational medicine research (protein and

biochemical markers, pharmacogenomics/pharmacogenetics biomarkers, pharmacodynamics) to its clients worldwide.

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