

A GLOBAL LEADER IN **BIOANALYSIS,** **PRECLINICAL AND CLINICAL** **RESEARCH SERVICES**

QPS is a Global Full Service CRO across USA, Europe and Asia offering Phase I-IV Drug Development Services



Pharmacology



Toxicology



DMPK



Bioanalysis



Translational
Medicine



Phase I Clinics



Regulatory &
Medical



Leukopak &
PBMC

FULL SERVICE GLOBAL CRO

- ▶ Pharmacology
- ▶ Toxicology
- ▶ DMPK
- ▶ Bioanalysis
- ▶ Translational Medicine
- ▶ Clinical Trial Conduct
- ▶ Clinical Research Services
- ▶ Cell Therapy and PBMC
- ▶ Clinical Trial Kits

CLINICAL RESEARCH OPERATIONS

- ▶ Project Management
- ▶ Clinical Program Management
- ▶ Clinical & Medical Monitoring
- ▶ Data Management & Biostatistics
- ▶ Medical Writing
- ▶ Quality Assurance
- ▶ Regulatory & Medical Affairs
- ▶ Safety & Pharmacovigilance
- ▶ Site Selection & Monitoring

THERAPEUTIC AREAS OF EXPERTISE

- | | | |
|-----------------|-------------------|------------------|
| ▶ Cardiology | ▶ MASH/NAFLD | ▶ Rheumatology |
| ▶ Diabetes | ▶ Neuropsychiatry | ▶ Vaccines |
| ▶ Dermatology | ▶ Obesity | ▶ Women's Health |
| ▶ Endocrinology | ▶ Oncology | |

Since 1995, QPS has provided discovery, preclinical, and clinical drug development services. An award winning leader focused on bioanalytics and clinical trials, QPS is known for proven quality standards, technical expertise, a flexible approach to research, client satisfaction and turnkey laboratories and facilities. For more information on QPS, visit www.qps.com, or email infobd@qps.com.

EARLY PHASE CLINICAL SITES

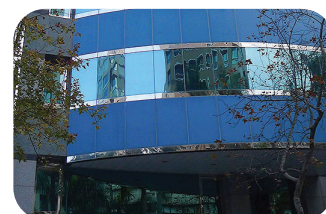
Miami

- ▶ 2 Clinical Units with 110 Phase I beds
- ▶ Over 3,000 studies completed, including POC, FIM, SAD, MAD, and specialty populations
- ▶ On-site board-certified physicians
- ▶ Database of over 35,000 potential study patients
- ▶ Excellent subject retention rates of over 95%
- ▶ Capabilities for over 65 consecutive inpatient nights
- ▶ 15 FDA inspections with no 483 reports
- ▶ Extensive experience in cardiology, diabetes, MASH/NAFLD, obesity, RA, vaccines, and more
- ▶ Certified PBMC lab and technicians onsite



Missouri

- ▶ 6 Clinical Units with 240 Phase I beds
- ▶ Over 2,000 studies completed including POC, FIM, SAD, MAD, and specialty populations
- ▶ On-site board-certified physicians
- ▶ Database of 46,000 potential study subjects
- ▶ On-site pharmacy, negative pressure room, and safety & central labs
- ▶ Leukopak and cell therapy facilities
- ▶ 31 FDA inspections with no 483 reports
- ▶ Clinical trial kit production and central lab kit logistics
- ▶ Certified PBMC lab and technicians onsite



Taiwan

- ▶ Full-service CRO offerings, serving global clients across Phase I-IV studies
- ▶ Over 365 studies completed, including POC, FIM, SAD, MAD, and specialty populations
- ▶ Clinical trial conduct, site selection and monitoring, project management, data management, biostatistics, report writing, medical and regulatory affairs, and IND packages
- ▶ Extensive experience in cardiology, diabetes, oncology, RA, and more
- ▶ 3 FDA inspections
- ▶ Toxicology and Pharmacology studies
- ▶ Bioanalysis and Sample Analysis

India

- ▶ 4 clinical units with 138 actively monitored beds
- ▶ Over 2,000 Phase I-IV studies
- ▶ BA/BE & PK/PD studies in healthy volunteers (biosimilars, DDI, dermal, inhalation, long acting injectables and PM women)
- ▶ PK/PD & clinical endpoint studies in patients (CNS, endocrinology, oncology, ophthalmology pulmonology)
- ▶ Database of over 23,000 potential study subjects
- ▶ 15 FDA, 1 ANVISA, 3 WHO, 2 MHRA, 1 EMA, 1 NPRA & 2 GCC successful inspections
- ▶ On-site sample collection/processing, pharmacy and safety lab
- ▶ Bioanalytical MD/MV and Sample Analysis