

A flexible approach to

# IND-enabling Preclinical Studies

**QPS IS A GLOBAL CRO WITH DIVERSE CAPABILITIES THAT OFFERS END TO END DRUG DEVELOPMENT SERVICES.**

Partnering with QPS for a well-conceived and executed IND-enabling preclinical program will provide you with a detailed assessment of your drug candidate and the most agile, flexible and timely pathway to filing an IND.

---

TIME IS OF THE ESSENCE IN DRUG DEVELOPMENT.  
CONTACT THE QPS BUSINESS DEVELOPMENT TEAM TODAY!

CALL +1 512 350 2827 EMAIL [infobd@qps.com](mailto:infobd@qps.com)



## Why perform your IND-enabling preclinical studies at QPS?

Before we begin executing your IND-enabling preclinical program, you will receive strategic review and advice on the design and execution of your ADME and pharmacology-toxicology studies:

- ▶ Your proposed non-clinical plan including proof of concept studies, pharmacology and ADME studies, and toxicology/safety program will be analyzed in depth
- ▶ Your proposed non-clinical plan will be reviewed to identify deficiencies and potential roadblocks and hurdles and whenever possible solutions identified
- ▶ Timelines for preclinical development of your overall and individual programs will be mapped out and preclinical development objectives and crucial milestones will be confirmed





## What are the Benefits of Working with QPS?

During execution of your IND-enabling program you will benefit from QPS's operational strengths, strong scientific/regulatory pre-IND/IND support, and drug development experience.

### Operational Strengths:

- ▶ ADME scientists and toxicologists with extensive industry and CRO experience allow for optimal planning and execution of ADME and pharmacology-toxicology studies
- ▶ State-of-the-art ADME, toxicology and bioanalytical facilities
- ▶ Rapid execution and completion of all preclinical studies required for IND submission
- ▶ All studies will be carefully monitored and every phase of the studies critically assessed for scientific rigor and quality
- ▶ Fast turnaround on high-quality non-clinical study reports
- ▶ Extensive experience in the preparation of ADME and pharmacology-toxicology sections of IND submissions
- ▶ An experienced program manager will be assigned to ensure rigorous program oversight

### Scientific/Regulatory Pre-IND/IND Support:

- ▶ ADME scientists and toxicologists with extensive industry and CRO experience allow for optimal planning and execution of ADME and pharmacology-toxicology studies
- ▶ Review and gap analysis of available data & preclinical development plans
- ▶ Advice on the design and timing of ADME, safety pharmacology, and toxicology studies
- ▶ Provide expert advice on ADME and pharmacology-toxicology issues associated with a broad range of therapeutic areas
- ▶ Rapid completion of the ADME and pharmacology-toxicology sections of the IND to enable client to file the IND in a timely manner



**AGILITY. FLEXIBILITY. SPEED**

# IND-enabling Preclinical Studies

**CALL** +1 512 350 2827  
**EMAIL** [infobd@qps.com](mailto:infobd@qps.com)

**QPS is a Global CRO**  
with locations around the world  
to serve the evolving needs of the  
Pharmaceutical and Biotech industries

