



# Non-Clinical Development

## OPTIMISING DRUG SAFETY EVALUATION FOR RAPID DECISION MAKING

**Non-clinical testing is conducted throughout all phases of drug development to assess the safety profile and pharmacokinetic/toxicokinetic (PK/TK) characteristics of drug candidates. If performed well, it can identify key ‘knock-out’ criteria early, maximizing the chances of success in subsequent clinical development.**

Strategies for the non-clinical development of candidate drugs follow general regulatory guidelines but are developed on a case-by-case basis according to the specifics of the drug. It is essential to design an optimal non-clinical development program that enables the medicinal product to be taken forward to the next phase in clinical development, or to product registration.



Venn has a team of non-clinical experts with broad experience who can guide you through all aspects of non-clinical development from lead candidate selection to market.

### Drug Safety Evaluation Strategy

Venn’s non-clinical consultancy services range from drug development planning to project outsourcing, monitoring and execution, reporting, writing expert summaries and reviews, collation of regulatory documentation and submissions.

Venn’s non-clinical experts can seamlessly integrate with your in-house experts and participate in multi-disciplinary development team(s) to facilitate accelerated drug development.

### Non-Clinical studies and related activities

Venn has broad experience in designing and monitoring non-clinical studies in all disciplines of toxicology, as well as in PK/TK and metabolism studies (ADME).

The professional management of a non-clinical project or program ensures that scientific and regulatory milestones are achieved, timelines are met, and budgets are respected.

Venn can take responsibility for execution of your non-clinical development program, by selecting the most suitable Contract Research Organization (CRO), discussing study designs, reviewing draft protocols, authorising final protocols as study monitor and reviewing draft study reports on your behalf.

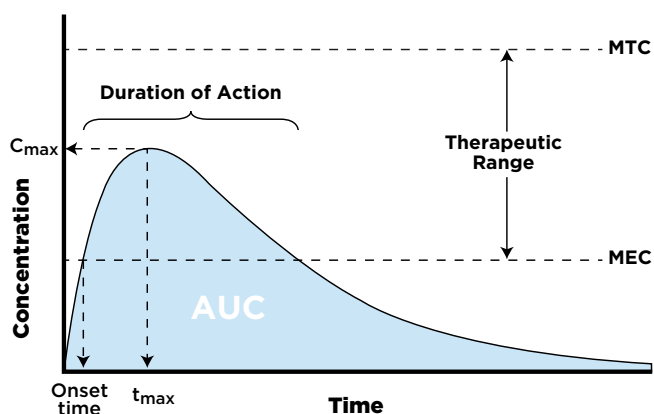
Our non-clinical expert will be the first point of contact with the study director at the CRO, enabling you to focus on other aspects of drug development.

## VENN NON-CLINICAL SERVICES

STRATEGIC	STUDIES	STUDY RELATED	REGULATORY DOCUMENTS
Non-Clinical Program Design & Management	Full range of Toxicity Studies (non GLP / GLP)	Study Design & Initiation	Investigator’s Brochure (IB)
Risk Analysis Risk Mitigation	ADME & PK / TK studies	CRO Selection & Management	Investigational New Drug (IND)
Consultancy, Advice, Support	Coordination & Monitoring of BA (Bioanalytical) studies	Execution & Coordination (onsite (insourcing) / off-site)	Clinical Trial Applications (CTA)
TPPs / DDPs Due diligence & GAP Analysis	PK / TK Evaluations	Analysis / Review & Reporting of Study Results	Expert Reports

## Non-Clinical PK/TK Evaluations

Venn offers Pharmacokinetic and Toxicokinetic (PK/TK) evaluation of bioanalytical data obtained from all types of non-clinical studies. The non-clinical department of Venn is accredited to perform these evaluations in accordance with the OECD principles of **Good Laboratory Practice (GLP)**.



The non-clinical department has broad experience in participating in multi-site studies, in collaboration with test facilities and test sites from all over the world. At request, our reports can be created in **SEND** format.

### EXPERIENCE OF VENN'S NON-CLINICAL DEVELOPMENT TEAM

Multidisciplinary - and Non-Clinical Projects (last 5 years): 35  
Regulatory Documents (last 5 years): 15  
PK/TK studies (last 5 years): 35

Broad Scientific Expertise & CRO Network

Small Molecules  
chemically synthesized products, synthetic oligonucleotides, synthetic oligopeptides, radiopharmaceuticals

Large Molecules  
Biological/biotechnological proteins and polypeptides, monoclonal antibodies, vaccines

Generics, Orphans Drugs, ADC's & Biosimilars

## Regulatory Documents

Venn can prepare all non-clinical documents for regulatory submissions for a variety of applications, such as non-clinical reports, non-clinical overviews, summaries and tables (INDs/CTAs), contributions to IBs and IMPDs. We can support you in discussions with the regulatory authorities by preparing briefing notes and Venn experts can accompany you to discuss non-clinical aspects of drug development with regulators.

## Why Venn's Non-Clinical Team

The combination of specific scientific (non-clinical) knowledge, commitment to your projects and the constant pursuit of quality excellence make Venn a reliable source for **Non-Clinical Consultancy Services** and for **Preparation of Regulatory Documentation** to support submission and a good choice for the outsourcing of **PK/TK Evaluation** according to GLP regulations, including writing of PK/TK study reports.



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