

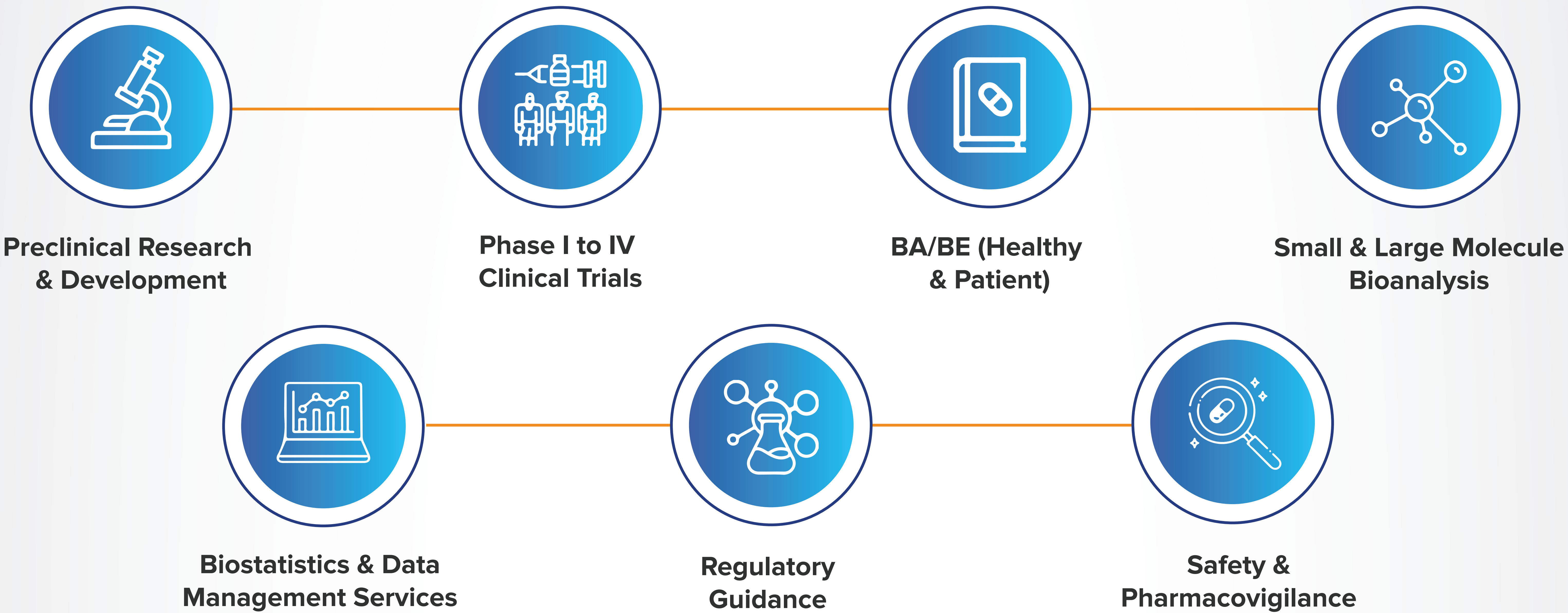
A Capable, Knowledgeable, and Reliable partner for your Drug-development journey



Veeda Group Overview

Veeda Clinical Research Limited (“Veeda”) together with its subsidiaries, Bioneds India Private Limited and Ingenuity Biosciences Private Limited (together referred to as the “Veeda Group”) offers a comprehensive portfolio of clinical, preclinical and bio/analytical services to support innovator, biosimilar and generic drug development programs of our global clientele. We are an independent, institutional investors owned, Board governed and professionally managed contract research group offering scientific leadership, global quality management systems and long term operational and financial stability through a continuing investment in our people, processes, systems, infrastructure and technology and a deep commitment to quality.

Our Integrated Drug-development Services



Veeda’s Vision

In an industry where innovation is increasingly multifaceted and collaborative, we aspire to be the research partner of choice for innovative (bio)pharmaceutical companies worldwide for their critical product development programs.

Veeda Clinical Research

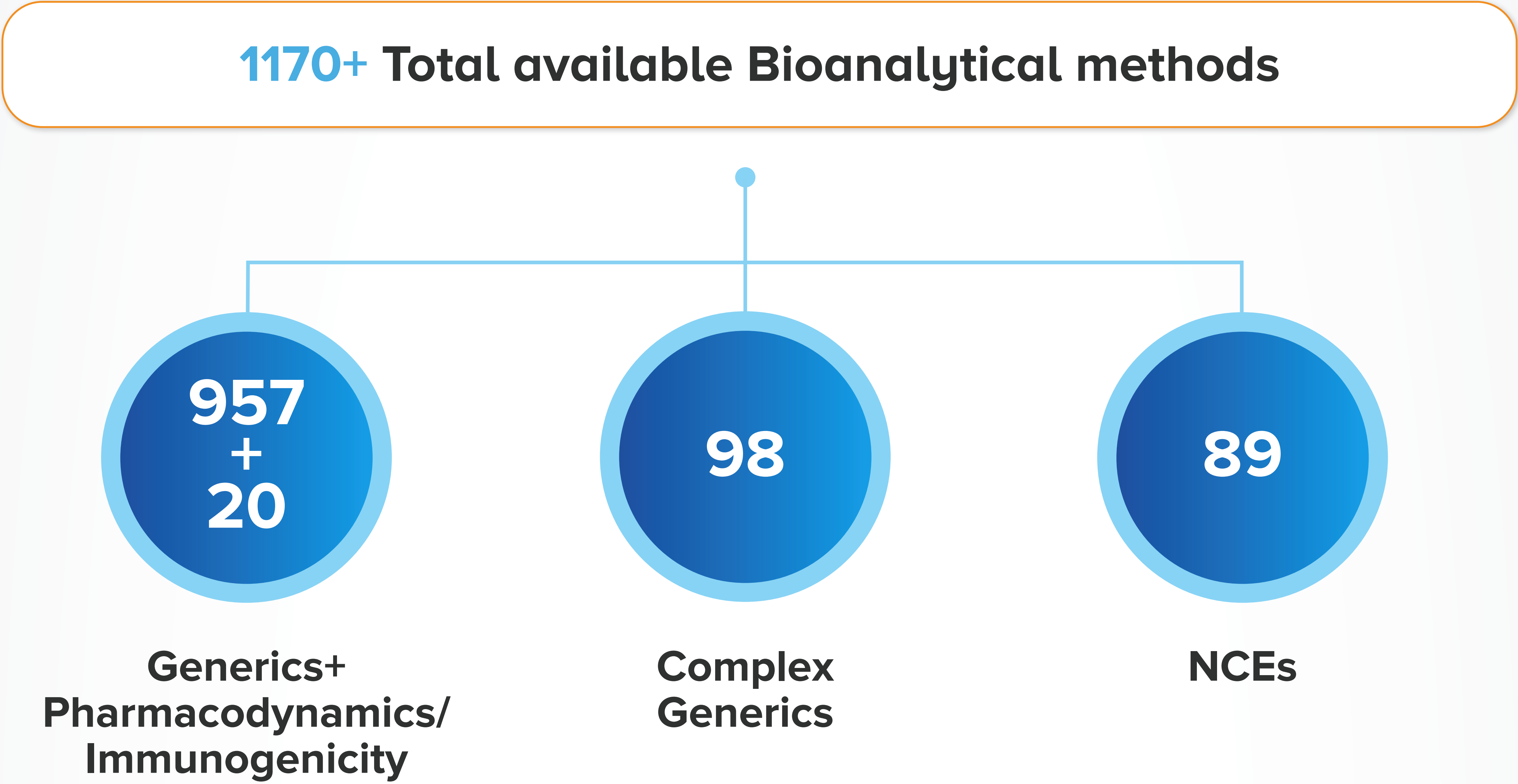
We provide access to Expertise & Knowledge that enables global (Bio)pharmaceutical companies to develop their new products

Our end to end services complement the research and development and marketing functions of global (Bio)pharmaceutical companies. Outsourcing these services to us enables our clients to move their molecules from preclinical development to clinical, and eventual commercialization in a timely and efficient manner

- Phase I Studies in Healthy Volunteers
- Phase I-IV Patient Trials
- Phase I/III Biosimilar Studies
- PK & Clinical Endpoint Studies
- Vaccine Trials
- Small & Large Molecule Bioanalysis

Strong Bioanalytical Capabilities to keep your Studies On Track

- Method development, validation & sample analysis for a wide range of drug substances
- Average processing capacity of 1,00,000 samples per month
- Central Bioanalytical Laboratory for global Phase II/ Phase III trials



Developed Complex Assays for: Endogenous molecules, Amino Acids, Hormones, Steroids, Inhalation formulation, Large molecules, Liposomal formulation, Exploratory studies (skin tissues, plasma protein binding experiment, chiral impurity estimation in the sample), Iron Sucrose, Peptides (small molecules).

Bionneeds: Globally Acclaimed Preclinical Contract Research Organization

With over 12 years of experience, Bionneeds is a leading Preclinical Contract Research Organization (CRO) providing Integrated Discovery, Development & Regulatory. Bionneeds has a state of the art facility with 200,000 sq ft built-up area in 5 acre campus in the outskirts of Bangalore.

Preclinical Services include

- > General Toxicity
- > Mutagenicity
- > DMPK
- > Immunotoxicology
- > Inhalation Toxicity
- > Eco Toxicity
- > Reproduction & Development Toxicity
- > Biological Tests
- > Physico Chemical Testing, Chemical/Drug Characterization

Ingenuity BioSciences

Ingenuity BioSciences is built on the complementary strengths to deliver Integrated Service Model for Drug Development, also bringing a strong synergy in offering a comprehensive bioanalytical solution to therapeutic and biosimilar development.

Non-Clinical Studies

- Structural Characterization**
Primary and higher order structures, Impurities, etc
- Functional Characterization**
Mechanism of action and cell-based signalling assays
- DMPK/ADME**
- Upstream & Downstream Capabilities**
Lab-scale manufacturing and purification, Cell-line characterization and qualification

Clinical Bioanalytical

- Vaccines**
Primary end-point analysis, Neutralizing antibodies, Serum Bactericidal Assay, PRNT/SNT
- Biologics/NBEs/Biosimilars**
Clinical Biomarkers, In-vitro Method Development & Validation, PK, PD, ADA, NAB assays Cytokine response

Our Regulatory Credentials

89 Successful Regulatory Audit till date

| | | | | | | | | |
|--------|---|----|----------|---|----|------|---|----|
| US FDA | → | 42 | WHO | → | 06 | AGES | → | 04 |
| MHRA | → | 04 | NPRA | → | 05 | MCC | → | 01 |
| ANVISA | → | 08 | Malaysia | | | DCGI | → | 18 |
| | | | ANSM | → | 01 | | | |



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