

NCE Bio analysis Capabilities

Veeda bioanalysis proficiency in handling new chemical entities (NCE) for different phase of clinical trials is backed by an experienced team of bio scientists having over 10 years of experience in handling NCE molecules. Veeda have 20 sensitive equipment's to support subpicogram level quantitation for the micro dosing studies to assess minimum effective dose.

The bio analytical team has handled various NCE molecules for different sponsors involving different phases of clinical trials and performed NCE analysis in targeted timeline to support:

- a) *Dose escalation – safety, PK & route*
- b) *Multiple dose studies to identify dose regime - frequency / route*
- c) *Formulation for effective bioavailability*
- d) *Populations based studies*

Veeda Advantage :

- Veeda has experience of about **36 validated NCE methods** for different sponsors involving **different biological matrices like human blood, serum, plasma, tissues and urine.**
- Validated method are having **dynamic range of over 2000 times linearity** to support SAD and MAD studies sample analysis.
- Veeda is committed to provide cutting edge methods to **Cross validation between sponsor lab and Veeda Lab** to adjudge method reproducibility and performance.

Veeda Advantage

- 100% data review by Bio-analytical Quality Monitors
- State of art Bio analytical Lab equipped with highend sensitive equipment's to achieve the required LLOQ.
- Trained Bio analysts to handle complex sample processing
- Proven regulatory track record with 11 USFDA, 5 European, 4 WHO & 11 ANVISA audits
- Bio-analytical Unit with more than 590 validated assays in its library of compounds including 36 NCE methods, 20 more under development.

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