veedacr.com

veeda edge

Optimize your complex Inhalation Clinical studies with one of india's most experienced CRO

Inhalation therapies are a group of respiratory treatments designed to help restore or improve breathing function in patients suffering from respiratory ailments. Nasal Drug Delivery is complex as it requires proper drug and device combinations for successful drug delivery at the site of action. Veeda Clinical Research Pvt. Ltd. is India's most experienced early phase clinical development CRO with a global outreach providing ethical Clinical Research Solutions. With a competent team of 650 experienced scientists and a stern focus on providing quality solutions, we are the ideal partner to accelerate your inhalation studies.

Requirements for Inhalation Studies

Scientific Expertise and through understanding of regulatory requirements in clinical development of inhalation products.

Temperature and Humidity monitored

2 Highly Compliant volunteers for Inhalation technique.

3

Clinical facility and IMP storage facilities.

Expertise in Conduct and Interpretation

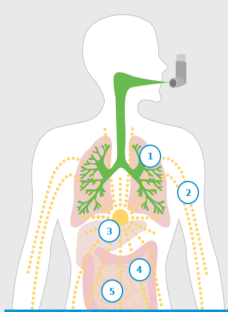
of Oxymetry and Spirometry.

Generic molecules for Inhalation studies

Fluticasone
 Ipratropium
 Salmeterol

Our Capabilities to ensure Successful Studies

- Vast experience of conducting pMDI & DPI studies on healthy volunteers.
- Team of professionals capable of managing highly complex inhalation studies.
- Team of professionals for training of volunteers and capable of performing & interpreting special tests such as Spirometry and Oxymetry.
- Dynamic database of highly compliant & well trained healthy male and female volunteers for complex studies such as PK/PD marker studies.
- Scientific Project Management Team sensitive to nuances of Inhalation studies.
- Ability to provide fast & cost effective solution for Inhalation studies.
- 100% data review by Quality Control / Quality Assurance.



Inhalation Chamber for Dosing

- Provides an uniform environment with relatively consistent temperature, humidity,air flow, oxygen,content and other major environmental factors for all respiratory dosing.
- 2. Eliminates any chance of cross contamination from one dosed subject to another during dosing procedure.
- 3. Better regulatory acceptance due to assured well-controlled dosing procedure.

Our Achievements

Successfully completed
6 Pharmacodynamic (PD)
& Clinical endpoint
studies for various
Inhalation Drugs.

Thorough Regulatory understanding and scientific expertise for Inhalation studies.

Data base of well trained volunteers.

India's most experienced early clinical development CRO.

State of the art to facility handle complex Inhalation Studies.

Proven regulatory track record with 20 USFDA (including 10 inspections at investigator sites), 5 European, 4 WHO & 5 ANVISA audits



For more information email: info@veedacr.com