veeda edge

Optimized practices to manage critical parameters for your Liposomal Doxorubicin Pharmacokinetic studies

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 **250 experienced scientists** and **a stern focus on providing quality solutions**, Veeda promises a well-planned conduct of Liposomal doxorubicin PK trials to ensure optimized trial outcomes by overcoming the challenges.

Challenges for Liposomal Doxorubicin PK trials



Standardization of **rate of infusion** is necessary to administer **precise dose** as per BSA without any **extravasation**.

Availability of suitable **Infrastructure at Clinical sites is essential** to effectively manage the critical controls and procedures to minimize variables.



Co-morbidities and duration of trial make **patient** retention and compliance difficult.

Differentiating between free and encapsulated doxorubicin is crucial during quantification in PK trials.



Our Capabilities to optimize your trial outcomes

- Data base of pre-screened experienced Investigator sites with requisite infrastructure.
- Validated bio-analytical method as per the US and the EU regulatory requirements.
- Identification of the critical controls that can affect the study outcomes and training the **site personnel to handle them effectively.**
- Complete cold chain management to ensure integrity of Investigational products and plasma samples.

Our Achievements

Recent success in completion of Renal Cell Carcinoma PK study within a very competitive timeline of 5 months	Fast recruitment rate even for rare indications like SCLC	Road Map for more than 25 molecules for Patient based Pharmacokinetic Studies and more than 15 molecules for Clinical End Point Studies	4 European,	State of the art Bio-analytical Unit with more than 340 validated assays in its library of compounds, 35 NCE methods and 20 more under development.
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