



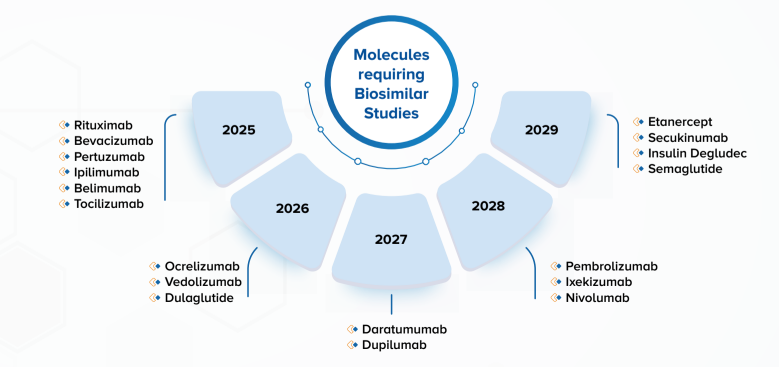
Injectable Drug Development Support

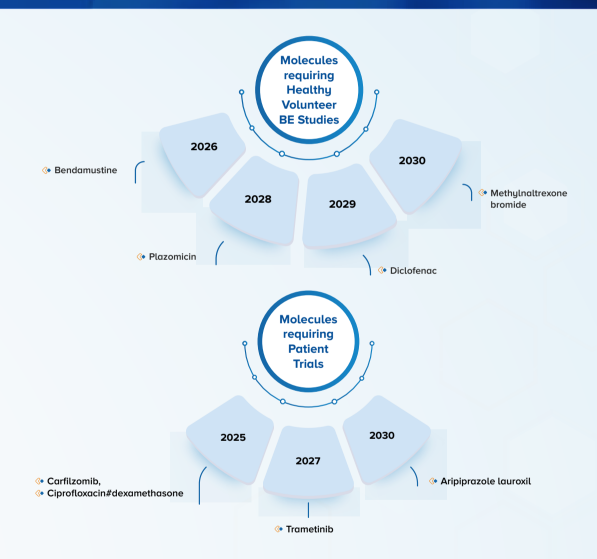
Complete Solutions for Injectable Drugs Nearing Patent Expiry: Ensuring Seamless Development & Market Readiness

Supporting You Across the Entire Drug Development Lifecycle

Preclinical	Biopharma	Clinical Studies	Bioanalytical
DiscoveryServices	Clinical Bioanalysis solution for Bio Therapeutics	Phase I Clinical Pharmacology studies for NCEs & NBEs	Method Development and Validation for NCEs and Generics
Preclinical	·		
Toxicology	Non-Clinical Characterization	Phase II- Phase III trials for NCEs & NBEs	
	Solutions – Discovery Biology, Bioprocess, and Analytical Characterization	Phase IV & PMS Studies	
◆ Bioanalysis		BE studies for Complex Injectable Drugs (both Healthy and Patient)	
	 Discovery Services Preclinical Toxicology Chemistry 	 Discovery Services Services Solution for Bio Therapeutics Preclinical Toxicology	 Discovery Services Services Solution for Bio Therapeutics Therapeutics Or NCEs & NBEs Phase I Clinical Pharmacology studies for NCEs & NBEs Or NCES & NBES Or

Injectable Drugs Coming off Patent in the Next Few Years





Injectable Drug Development comes with Unique Set of Challenges







Injectable drugs need specific storage for stability



Regulatory Approval Processes: The approval process can be lengthu and complex requiring comprehensive data on safety, efficacy, and manufacturing practices

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Trials requiring dose adjustments based on patient



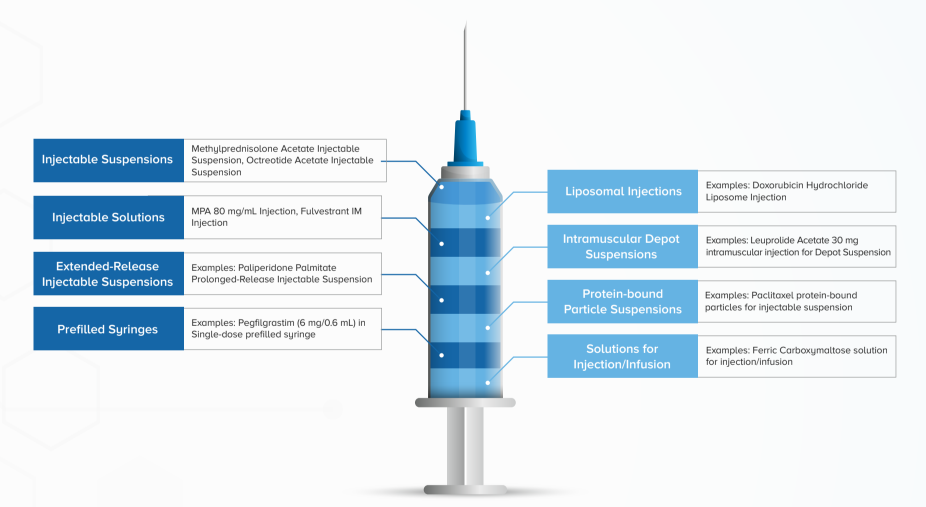


Veeda's Expertise in Injectable Studies

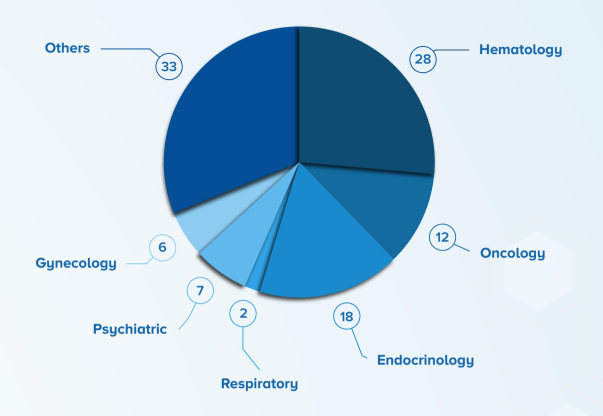
Comprehensive Bioequivalence Research: 106 Healthy Volunteers and **Patient Studies**



Wide Experience Across Different Complex Formulation



Trusted Companion for Injectable Drug Development Across Diverse Therapy Areas



Bioanalytical Method **Validation**

Experts in utilizing advanced techniques such as mass spectrometry and HPLC for precise and thorough analysis.

Successfully supported a range of injectable studies through a range of validated methods via LC-MS/MS, like injectable suspensions (e.g., Octreotide Acetate), proteinbound particle suspensions (e.g., paclitaxel), liposomal injections (e.g., doxorubicin & Amphotericin B), subcutaneous injections (e.g., Phytonadione) and many more.

Advanced Analytical Techniques and Clinical Bioanalysis for Injectable Biosimilars



Bioanalytical Expertise

Specializing in clinical and bioanalytical evaluation, with advanced techniques such as ELISA and LC-MS/MS for immunogenicity assessment.



Peptides Expertise

Advanced analytical techniques for peptide characterization, peptide PK/PD studies, and peptide mapping to ensure a comprehensive analysis for various peptides like Insulin, Desmopressin, Leuprolide, and Octreotide.



Biosimilars Experience

Extensive experience in the development and evaluation of biosimilars, including ADL-018 (XOLAIR), a recombinant DNA-derived monoclonal antibody.



Experience with Different Molecules

Teriparatide, Tocilizumab, Romiplostin, FSH (Follicle Stimulating Hormone), and C-Peptide.





