

Voluntarily

## 16<sup>th</sup> Munich Workshop on VICH GCP and Veterinary Clinical Studies

April 4th -5th, 2019

Maritim Hotel München, Goethestr. 7, 80336 München/ Germany

## Day 1 – April 4<sup>th</sup>, Thursday

Time	Topic	Presenter	
12:00 – 12:30	Registration and Snacks		
12:30 – 13:00	Welcome and introduction to veterinary GCP	Klaus Hellmann	
		Klifovet AG,	
		Germany	
13:00 – 13:30	Strategic planning – How to get on the right track	Hedi Goerg	
	What is strategic planning?	Klifovet AG,	
	<ul> <li>What is involved with strategic planning in the context of clinical development?</li> </ul>	Germany	
	How to adapt your strategy?		
13:30 – 14:15	Dose finding and confirmation – Defining suitable efficacy parameter	Klaus Hellmann Klifovet AG,	
	How to define suitable efficacy parameters	Germany	
	Statistical significance versus clinical relevance		
	Interpretation of results		
	Consequences for SPC		
14:15 – 14:45	Coffee Break		
14:45 – 15:15	Considerations for studies under field conditions	Claudia Schneider	
	General requirements	Klifovet AG,	
	Study design	Germany	
	Formal requirements to the study protocol		
15:15 – 16:00	The regulatory view in the assessment of clinical	Gesine Hahn, BVL,	
	efficacy	Germany	
	What needs to be covered?	·	
	What are the potential pitfalls?		
	The importance of efficacy in the benefit-risk assessment		
16:00 – 17:30	Workshop: Design of clinical studies – Preparing a	Claudia Schneider	
	study outline	& Lena Naderer	
		Klifovet AG, Germany	

From 19:00

Networking Dinner



## Day 2 – April 5<sup>th</sup>, Friday

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Time	Topic	Presenter	
08:30 – 09:15	Responsibilities in clinical studies	Gabriele Braun	
	Sponsor, Monitor, Investigator	Klifovet AG, Germany	
09:15 – 09:45	Setting up clinical studies in the field	Claudia Schneider	
	Investigator selection	& Kerstin Adler	
	Patient recruitment and follow-up	Klifovet AG, Germany	
09:45 – 10:45	Workshop: Monitoring of clinical studies – A case study	José Matallo Klifovet AG, Germany	
10:45 – 11:15	Coffee Break		
11:15 – 11:45	Clinical supplies requirements and obtaining regulatory	Klaus Hellmann	
	approval	Klifovet AG, Germany	
11:45 – 12:15	Assuring quality in clinical studies	Claudia Laskowski	
	Background – Quality management of clinical studies	Klifovet AG,	
	Elements for Quality	Germany	
	QA vs. QC		
	Quality assurance and auditing		
	How to prepare a study site for an inspection		
12:15 – 13:15	Lunch break		
13:15 – 14:15	Data Management	Dejan Cvejić	
	GCP requirements	Klifovet AG,	
	Electronic vs. Paper data capturing and processing	Germany	
	Data analysis and reporting		
14:15 – 14:45	Coffee break		
14:45 – 15:15	Practical statistics planning and assessment	Jose Matallo	
	Population and sample	Klifovet AG,	
	Distribution and probability	Germany	
	Types of data and their evaluation		
	Hypotheses and errors		
	Confidence intervals and sample sizes		
15:15 – 15:45	Benefit- risk balance: the assessment of authorities	Gesine Hahn,	
		BVL, Germany	
15:45 – 16:00	Closing remarks	Klaus Hellmann	