

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

April 27<sup>th</sup> – 28<sup>th</sup>, 2017

NH Hotel Deutscher Kaiser, Arnulfstr. 2, 80335 München/ Germany

## Day 1 – April 27<sup>th</sup>, Thursday

Time	Торіс	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	Regina Wolf
	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 view of regulatory assessor in clinical efficacy	Laure Baduel
	studies	ANSES,
	What needs to be covered	France
	• Miletale and the metantial withfalls in classican?	
	<ul> <li>Which are the potential pitfalls in dossiers?</li> </ul>	
	<ul> <li>Which are the potential pittalis in dossiers?</li> <li>The importance of efficacy on benefit/ risk assessment</li> </ul>	
16:00 - 17:15		Miriam Haas
16:00 - 17:15	The importance of efficacy on benefit/ risk assessment	Miriam Haas Klifovet AG,
16:00 - 17:15	<ul> <li>The importance of efficacy on benefit/ risk assessment</li> <li>Workshop: Design of clinical studies – Preparing a</li> </ul>	



## Day 2 – April 28<sup>th</sup>, Friday

Time	Topic	Presenter
08:30-09:15	Responsibilities in clinical studies	Gabriele Braun
	Sponsor, Monitor, Investigator	Klifovet AG,
09:15 - 09:45	Satting up clinical studies in the field	Germany Miriam Haas
09.15-09.45	Setting up clinical studies in the field	
	Investigator selection	Klifovet AG, Germany
	Patient recriutment and follow-up	,
09:45 – 10:45	Workshop: Monitoring of clinical studies – A case study	José Matallo
		Klifovet AG,
		Germany
$\frac{10:45 - 11:15}{11:15 - 11:45}$	Coffee Break Clinical supplies requirements and obtaining regulatory	Klaus Hellmann
11.15 11.45	approval	
		Klifovet AG, Germany
11:45 - 12:15	Assuring quality in clinical studies	Claudia Laskowski
	<ul> <li>Background – Quality management of clinical studies</li> </ul>	Klifovet AG,
	Elements for quality	Germany
	<ul> <li>QA vs. QC</li> </ul>	,
	Quality assurance and auditing	
	<ul> <li>How to prepare a study site for an inspection</li> </ul>	
10.15 10.15		
12:15 - 13:15 13:15 - 14:15	Lunch break Practical statistics planning and assessment	Hannes Buchner
10.10 11.10	<ul> <li>Population and sample</li> </ul>	Staburo Statistical
	Distribution and probability	Consulting GmbH,
	<ul> <li>Types of data and their evaluation</li> </ul>	Germany
	Hypotheses and errors	
	<ul> <li>Hypotheses and errors</li> <li>Confidence intervals and sample sizes</li> </ul>	
	Confidence intervals and sample sizes	
14:15 - 14:45	Confidence intervals and sample sizes     Coffee break	Doion Cuoilá
14:15 – 14:45 14:45 – 15:15	<ul> <li>Confidence intervals and sample sizes</li> <li>Coffee break</li> <li>Data Management</li> </ul>	Dejan Cvejić
	<ul> <li>Confidence intervals and sample sizes</li> <li>Coffee break</li> <li>Data Management</li> <li>GCP requirements</li> </ul>	Klifovet AG,
	<ul> <li>Confidence intervals and sample sizes</li> <li>Coffee break</li> <li>Data Management</li> <li>GCP requirements</li> <li>Electronic vs. Paper data capturing and processing</li> </ul>	
14:45 – 15:15	<ul> <li>Confidence intervals and sample sizes</li> <li>Coffee break</li> <li>Data Management</li> <li>GCP requirements</li> <li>Electronic vs. Paper data capturing and processing</li> <li>Data analysis and reporting</li> </ul>	Klifovet AG, Germany
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