

14th Munich Workshop on VICH GCP and Veterinary Clinical Studies

April 27th – 28th, 2017

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

Day 1 – April 27th, 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 <ul style="list-style-type: none"> • What is strategic planning? • What is involved with strategic planning in the context of clinical development? • How to adapt your strategy? 	Regina Wolf Klifovet AG, Germany
13:30 – 14:15	Dose finding and confirmation – Defining suitable efficacy parameter <ul style="list-style-type: none"> • How to define suitable efficacy parameters • Statistical significance versus clinical relevance • Interpretation of results • Consequences for SPC 	Klaus Hellmann Klifovet AG, Germany
14:15 – 14:45	Coffee Break	
14:45 – 15:15	Considerations for studies under field conditions <ul style="list-style-type: none"> • General requirements • Study design • Formal requirements to the study protocol 	Claudia Schneider Klifovet AG, Germany
15:15 – 16:00	The view of regulatory assessor in clinical efficacy studies <ul style="list-style-type: none"> • What needs to be covered • Which are the potential pitfalls in dossiers? • The importance of efficacy on benefit/ risk assessment 	Laure Baduel ANSES, France
16:00 – 17:15	Workshop: Design of clinical studies – Preparing a study outline	Miriam Haas Klifovet AG, Germany
From 19:00	Networking Dinner	Voluntarily

Day 2 – April 28th, Friday

Time	Topic	Presenter
08:30 – 09:15	Responsibilities in clinical studies <ul style="list-style-type: none"> • Sponsor, Monitor, Investigator 	Gabriele Braun Klifovet AG, Germany
09:15 – 09:45	Setting up clinical studies in the field <ul style="list-style-type: none"> • Investigator selection • Patient recruitment and follow-up 	Miriam Haas 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 approval	Klaus Hellmann Klifovet AG, Germany
11:45 – 12:15	Assuring quality in clinical studies <ul style="list-style-type: none"> • Background – Quality management of clinical studies • Elements for quality • QA vs. QC • Quality assurance and auditing • How to prepare a study site for an inspection 	Claudia Laskowski Klifovet AG, Germany
12:15 – 13:15	Lunch break	
13:15 – 14:15	Practical statistics planning and assessment <ul style="list-style-type: none"> • Population and sample • Distribution and probability • Types of data and their evaluation • Hypotheses and errors • Confidence intervals and sample sizes 	Hannes Buchner Staburo Statistical Consulting GmbH, Germany
14:15 – 14:45	Coffee break	
14:45 – 15:15	Data Management <ul style="list-style-type: none"> • GCP requirements • Electronic vs. Paper data capturing and processing • Data analysis and reporting 	Dejan Cvejić Klifovet AG, Germany
15:15 – 15:45	Benefit/ Risk assessment – The approach of the regulatory authorities	Laure Baduel ANSES, France
15:45 – 16:00	Closing remarks	Klaus Hellmann Klifovet AG, Germany