

CRED Navigating European Regulatory Procedures 15-16 March 2022

Day One - 15 March 2022

Time (GMT)	Session	Presenter
08.30	Registration and coffee	
09.00	Chairman's Introduction	Robert Clay Highbury Regulatory Science Limited
09.05	Case study introduction	Matthew Sardo
	 Delegates will be divided into groups for afternoon case study and provided with material. 	Sardo Trading Limited
09.10	Overview of the European Regulations	Kora Doorduyn- van der Stoep Medicines Evaluation Board (MEB)
	 A brief outline of the Centralised and Mutual Recognition/Decentralised procedures 	
	 The role of the regulatory bodies, i.e. EMA, CHMP, COMP, PDCO, SAGs, EU Commission, CMDh, National Authorities, ATMP, PRAC, EMA re-structure 	
	Orphan Drug Regulations	
	Paediatric Regulations	
	PRIME and similar initiatives	
	Considerations for SMEs	
	The role of patients	
10.10	Developing your Global Filing Strategy	Robert Clay
	 Influencing factors, e.g. product type, strength of data, internal & external initiatives (include in learning objectives) 	Highbury Regulatory Science Limited
	Scientific advice – where and how to obtain it	
	Utilising internal and external expertise	
	Integrating filing strategy with commercial objectives	
	 Choosing a procedure – key factors (default procedure unless otherwise indicated/company preference?) 	
	Orphan Drug Considerations	
	Paediatric Study Requirements	
	Accelerated approval programs	
	Briefly mention pharmacoeconomics and HTA requirements	
	Other initiatives (e.g. PRIME)	
11.00	Networking session	
11.30	Tea/ Coffee break	
12.00	Developing Strategy for Filing Non-NCEs	Eesha Vyas
	Bibliographic/ established use	GSK



Time (GMT)	Session	Presenter
	 Abridged Line extensions/generics/hybrids Biosimilars Borderline and combination products Herbals 	
13.00	Lunch	
14.00	 Decentralised and Mutual Recognition Procedures Background to mutual recognition concept Key features and differences of MRP/DCP Role of the Reference Member State and booking your RMS slot DCP procedure timetable Organising your company Interacting with the Authorities during the procedure Common problems and pitfalls in DCP Repeat-use MRP The role of CMDh Appealing a negative decision - referral and arbitration Common problems and pitfalls in MRP 	Stephen Smith MESAVS Regulatory Solutions Ltd
15.00	Networking session	
15.30	 Finalising your European Regulatory Strategy Scheduling the submission Access to procedures Commercial considerations Tradename issues Pre-submission meetings Timelines Conditional approvals/exceptional circumstance 	Peter Embley Bionical Emas
16.30	Case Study (Coffee/tea to be taken in case study groups)	Matthew Sardo
17.15	Case Study Feedback Presentation of results and discussion	
17.45	Review of the day	Robert Clay Highbury Regulatory Science Limited
18.00	Close of Day One	



Day Two - 16 March 2022

Time	Session	Presenter
08.30	Introduction	Robert Clay
	Review of day 1	Highbury Regulatory
	Objectives for day 2	Science Limited
08.40	Key EU Regional Considerations	Philip Senior
	Electronic dossier format (eCTD)	Pfizer Limited
	Labelling and leaflets including readability testing	
	Translations	
	Pharmacovigilance Site Master File and Risk Management Plan	
	Environmental Risk Assessment	
	Paediatric Investigational Plans	
	 Manufacturing aspects – site inspections, QC testing and QP release 	
09.40	Networking session	
10.00	Centralised Procedure - The Details	Claire Levee
	Preparing for a CP (including eCTD compilation)	Regulatory Consultant
	Managing the procedure	
	Interacting with the EMEA/(co-)rapporteur/PRAC	
	The role of the CHMP	
	The Commission decision making process	
	Re-examination of an Opinion	
	Organising your company	
	Common problems and pitfalls	
10.30	Tea/ coffee break	
11.00	Agency perspective on strategic and practical considerations – how to work effectively	Kora Doorduyn-van der Stoep
	 Developing and optimising your strategic plan – how to maximise the chances of an early approval 	Medicines Evaluation Board (MEB)
	Scientific advice and its impact on a successful application	
	 Validation, responses to questions and end of procedure – best practice advice 	
	 Typical dossier deficiencies and how to avoid them (no. of dossiers that go through first time and if not, why not) 	
	Common pitfalls (both practical and strategic)	
	 Real-life procedure timings – where time can be lost in a procedure 	
	Talking to regulatory agencies	
	Optimising appeals and hearings	
	Scientific Advice	



Time	Session	Presenter
12.00	New MAAs in the Centralised Procedure – Theory versus Reality • Understanding the potential for deviating from the rule book • Impact of PRAC • Impact of CAT • Impact of ODD • What are the potential outcomes at D181? • Thinking strategically Learning to manage internal expectations	Paul Nitschmann Intercept Pharmaceuticals

13.00	Lunch	
13.45	Networking session	
14.15	Case Study	Matthew Sardo
		Sardo Trading Limited
16.00	Case Study Feedback	
	Presentation of results and discussion	
16.30	Review of the day	Robert Clay
		Highbury Regulatory Science Limited

16.45 Close of Meeting

Delegates will be encouraged to ask questions throughout the day so as to ensure the meeting is as interactive as possible. There will be an interactive session with each speaker, at the end of each talk to consolidate the key points of each talk, and to allow the delegate to interact with all of the speakers.