Module 5: Regulatory Control of Clinical Operations 17th – 19th July 2024 **Location:** TOPRA Office, 6th Floor, Harbour Exchange, London, E14 9GE and Online



Module Leader(s): Eva Kopecna and TBC

Date: Wednesday 17th July 2024

Time	Activity	Speaker
13.00 - 13.15	Welcome & Introduction To Module 5	Chairperson TBC
13.15 - 14.15	Lecture 1: GCP - The platform for clinical research. Objective History Quality & Compliance Audits and Inspections	Eva Kopecna Acino International (F2F)
14.15 - 15.15	Lecture 2: GCP Inspections – Agency Experience	Rachel Mead MHRA (video)
15.15 - 15.45	Refreshment break	
15.45 - 17.00	Lecture 3: Progression of clinical trials in relation to the toxicology programme Purpose of toxicology studies in support clinical development Requirements for different stages of clinical development Requirements for different types of medical products (e.g. chronic, acute, delivery system)	Simon Craige EdGe Toxicology Consulting (F2F)



Date: Thursday 18th July 2024

Time	Activity	Speaker
09.00 - 09.15	Introduction	Chairperson: TBC
09.15 - 10.15	Lecture 4: Regulatory requirements for clinical trials - EU Current Legislation Requirements The Clinical Trial Regulation and its Impact CTA applications / Medical Devices Ethics Committee Submissions Reporting of clinical trials.	Pierre Omnes Transperfect Life Sciences
	Industry experience	
10.15 - 10.45	Refreshment Break	
10.45 - 11.45	Lecture 4: Regulatory requirements for clinical trials – EU Cont'd	Pierre Omnes Transperfect Life
11.45 - 12.45	Lecture 5: The successful IMPD Contents of the IMPD Pitfalls Updates	Graham Bell ICON
12.45 - 13.45	Lunch	

13.45 - 14.30	Lecture 6: GMP, authorisation and importation requirements, clinical trial supplies management 2003/94/EC GMP Directive Manufacturing and importation requirements including the role of the QP Regulatory framework under which clinical	Venkat Sunkara ICON
	supplies are managed Planning of a clinical supplies programme Issues relating to manufacture, packaging and distribution of clinical supplies	

14.30 -	- 15.15	Lecture 7: Pharmacovigilance & adverse event reporting in clinical trials Adverse Event reporting -General requirements and definitions Safety reporting in the EU Pharmacovigilance. Future Proposals	Beatrice Panico MHRA (remote)
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15.15 – 15.45 Refreshment Break

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15.45 – 16.00 Introduction to Case Study 1 TBC	
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16.00 - 17.00	Case Study 1: Initiation of a Clinical Trial
17.00 - 17.30	Presentation of feedback from Case Study 1

Date: Friday 19th July 2024

Time	Activity	Speaker
08.45 - 08.50	Introduction	Chairperson Eva Kopecna Acino International
08.50 - 09.50	Lecture 8: Legal aspects of clinical trials Product Information and Informed Consent Data Protection. Enforcement and Sanctions Liability. Clinical Study Contracts	Adela Williams Arnold and Porter (remote)
09.50 - 10.30	Refreshment Break	
10.30 - 11.30	Lecture 9: Regulatory requirements for clinical trials - Japan Legislation requirements Clinical Trial Notifications Japanese Regulatory Authorities Consultations with the Authorities Clinical Trials in Japan & GCP	Mohamed Oubihi Yakumed
11.30 - 12.30	Lecture 10: Regulatory requirements for clinical trials – US and Canada Legislation requirements CTA and IND applications & maintenance Ethics Committee GCP & Enforcement	Sarah Roberts PRAHS
12.30 - 13.30	Lunch	
13.30 - 13.45	Introduction to Case study 2: GCP and Clinical Development Programme	Eva Kopecna Acino International
13.45 - 14.45	Case Study 2: GCP and Clinical Development Programme	
14.45 - 15.15	Presentation of feedback from Case Study 2	

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