

## CRED Compiling Successful Clinical Trial Applications 7-8 February 2024, Prague

## **DAY 1: Compiling Successful Clinical Trial applications**

Chairperson: Shaila Choi, Seagen International GmbH

Time	Session/ Activity	Presenter
09:00	Registration and Coffee	
09:15	Welcome from TOPRA	TOPRA
09:20	Introduction to Clinical trials and the next 2 days	<b>Shaila Choi</b> Seagen International GmbH
09:45	<ul> <li>Essential Documents</li> <li>ICH/GCP</li> <li>IB</li> <li>Protocol</li> <li>IMPD</li> <li>Risk:Benefit</li> <li>Comparators</li> <li>RSI</li> </ul>	<b>Stephen Thompson</b> S Cubed
11:15	Tea and Coffee	
11:45	<ul> <li>Overview of the EU-CTR</li> <li>Initiation of clinical trials</li> <li>Regulation 536/2014</li> <li>Portal and Database</li> </ul>	Pierre Omnes TransPerfect  Matthew Scullion
	<ul> <li>Practical considerations</li> <li>Experience</li> </ul>	Janssen
13:00	Lunch	
14:00	<ul> <li>How to maintain your CTA: CTD</li> <li>Maintenance of the CTA: <ul> <li>How to navigate the bumps and pitfalls;</li> <li>Amendments;</li> <li>Temporary halts;</li> <li>Safety reporting;</li> <li>End of trial activities;</li> <li>Reporting obligations;</li> <li>Archiving/TMF maintenance</li> <li>Clinical trial Transparency and Disclosure and maintenance of registries.</li> </ul> </li> </ul>	<b>Stephen Thompson</b> S Cubed
14:45	<ul> <li>How to maintain your CTA: CTR 536/2014</li> <li>Maintenance; Modifications, Temp Halts, Safety reporting, EoT activities, reporting obligations, Archiving,</li> <li>Transparency &amp; Disclosure</li> <li>Transition considerations: CTD to CTR</li> </ul>	Pierre Omnes Transperfect  Matthew Scullion Janssen
15:45	Tea and Coffee	
16:15	CTR Demo	Pierre Omnes Transperfect
17:30	Close of meeting - Day 1	



## **DAY 2: Global considerations for Clinical Trial Applications**

Chairperson: Stephen Thompson, S Cubed

Time	Session/ Activity	Presenter
09:15	Welcome and Introduction to the day	<b>Stephen Thompson</b> S Cubed
09:20	<ul> <li>GCP &amp; inspections</li> <li>GCP and what the RA professional involved in Clinical Trial Management needs to know</li> <li>Overview and types of GCP Inspections</li> <li>How the RA professional needs to prepare for an Inspection</li> <li>What to expect in the Inspection</li> <li>TMF</li> </ul>	<b>Bob Ibbotson</b> Shionogi BV
10:30	Tea and Coffee break	
11:00	RoW CTs:  UK combined review process FDA SwissMedic LATAM APAC	<b>Stephen Thompson</b> S Cubed
12:30	Lunch	
13:30	Global Regulatory Lead: Product development & Strategic considerations  • Study design • Choice and provision of comparators, populations, SOC • Considerations for ATMPs and GMOs • How regulatory interacts with clinical in clinical development Discussion led by Stephen Thompson – Global Development requires global data  Discussion led by Pierre Omnes– Post study treatment.  What happens to patients following completion of the study? - Compassionate use/named patient supply.  Tea and coffee break	<b>Matthew Scullion</b> Janssen
14:30 15:00	Agency's perspective on CTAs	Eva Hrušková
15:00	<ul> <li>Hot Topics and Common Issues with CTAs</li> <li>CTR updates</li> </ul>	SUKL
16:00	Closing summary of the day	<b>Stephen Thompson</b> S Cubed
16.30	Close the meeting	