

For the full course descriptions and the latest updates visit our website:

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	Course Name	Type	Location	Jan	Feb	Mar	Apr	May	Jun	Jul	Sep	Oct	Nov	Dec	
<b>T</b>	<b>Clinical Operations Courses</b>														
	Introduction to Clinical Research	E													
	Orienting your Career in Clinical Research	C	Brussels									02			09
		C	Leiden		18										
	Foundational ICH-Good Clinical Practice (GCP) E6 (R2) Training	C	Brussels			25			13		01	03			10
		C	Milan								09				
		C	Leiden		19										
		E													
	ICH-Good Clinical Practice Refresher	E													
	ICH-GCP E6 (R2) Addendum 2016														
	ICH-Good Clinical Practice Refresher + Complementary Module GCP Refresher for Clinical Operations Staff*	E													
	ICH-Good Clinical Practice Training for Investigators	E													
	Clinical Research Training for Clinical Trials Assistants (CTAs)	C	Brussels											04-05	
	Clinical Research Training for Junior Clinical Research Associates (CRAs)	C	Brussels			26-27			14-15		02-03	04-05			11-12
		C	Milan								10-11				
		C	London							25-26					
		C	Leiden		20-21										
		C	Munich				10-11								
	Clinical Research Training for Senior Clinical Research Associates (CRAs)	B	Brussels		25					17		23			
		B	Milan						27						
	Risk Based Monitoring	B	Brussels		28							27			
	Introduction to Clinical Research with Medical Devices	C	Brussels						06						
	ISO 14155 Training	C	Brussels						07						
	Running Medical Device Trials	C	Brussels						09-10				21-22		
	<b>Regulatory Courses</b>														
	Basics on Regulatory Requirements in Clinical Research	C	Brussels									02			09
C		Leiden		18											
European Legislation for Clinical Research - Implementation in Belgium	C	Brussels								26		28			

\*These trainings are only available once ICH-GCP refresher is completed.

B= Blended (eLearning+Classroom), C= Classroom, E= eLearning, W= Webinar

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Medical Device Regulations	C	Brussels					08				23		
The Belgian Clinical Trials Law of 2017: A Clear View on Rules	W				18		27		08		24		16
The European Clinical Trial Regulations 536/02014 - A Clear Outline	W		24			08		19			14		
European Legislation of Clinical Trials with Medicinal Products	C	Brussels								12		25	
	E												
ICH-GCP Refresher + Complementary Module GCP Refresher for Regulatory Staff*	E												
Comparing EU with US Legislative Requirements of Clinical Trials	E												
Local Clinical Trial Regulations	E	Austria, Belgium, Czech Republic, Denmark, Finland, France, Germany, Hungary, Italy, Norway, Poland, Russia, Spain, Sweden, The Netherlands and UK <b>(More EU Countries to be added in 2019)</b>											
<b>QA Related Courses</b>													
Introductory Course on Auditing Investigator Sites	C	Brussels				02-03					23-24		
Introduction to Inspection Readiness	C	Brussels				05				25			
Introduction to System Audits for Clinical Auditors	C	Brussels				02							
Computer System Validation for Clinical Operations	C	Brussels		07-08									
<b>T</b> <b>NEW</b> Writing Audit Reports	C	Brussels				01							
<b>NEW</b> Risk Based Approach to Clinical Audits	C	Brussels				03							
<b>NEW</b> Auditing Clinical Development Documents	C	Brussels				04							
<b>NEW</b> Communication and Appreciative Auditing	C	Brussels				04							
<b>NEW</b> Clinical Service Provider Audits	C	Brussels				04							
<b>NEW</b> Sponsor Co-Monitoring	C	Brussels						20					
<b>Clinical Research Related Courses</b>													
Legal Basics for Clinical Study Contracts	C	Brussels			15						11		
Data Protection in Clinical Research and GDPR in action	C	Brussels	28							09			
Bridging Pre-Clinical and Clinical Research	C	Brussels		21-22									
Liability & Insurance in Clinical Research in Belgium	C	Brussels	28							09			
Introduction to Oncology for Clinical Researchers	C	Brussels			28-29								
Introduction to Statistics	C	Brussels						06-07					
Pharmacovigilance in Clinical Trials	C	Brussels						12					
Clinical Development of a Vaccine	C	Brussels										25-26	
Paediatric Clinical Development	C	Brussels										27-28	

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	Good Manufacturing Practice (GMP) in relation to GCP	E												
	ICH-GCP Refresher + Complementary Module GCP Refresher for Biometrics Staff*	E												
M	Clinical Project Management	C	Brussels			11-12			03-04			07-08		05-06
		C	London						27-28					
		C	Munich					27-28						
	Risk Management in Clinical Research	B	Brussels			14						10		
	Advanced Clinical Project Management	C	Brussels					06-07					18-19	
	Time Management	C	Brussels		26						24			
	CRO Management	C	Brussels					09-10					21-22	
	Train the Trainer	C	Brussels			14								
	MS Project Basics for Clinical Project Managers	W			19,21	02,04			18,20	02,04		01,03 15,17		
L	People Management	C	Brussels			13						09		
	Leading in a Solution Focused Way	C	Brussels					08					20	
	Line Management Essentials	C	Brussels									14-15		

## Other courses available:

Besides our planned courses, we have other courses also available for your training needs. Contact us if any of them are of interest to you and we will work on scheduling them:

- Intercultural Communication
- Change Management
- Introduction to Clinical Data Management for Clinical Researchers
- IMP Manufacturing & Management
- Effective Medical Writing & Data Presentation
- The ECG in Clinical Research
- Laboratory Testing in Clinical Research

## Contact us for more information:

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