

GREAT!

YOU'VE JUST FOUND

**THE PERFECT
CLINICAL RESEARCH
LEARNING & DEVELOPMENT
SOLUTION**



EUROPEAN CENTRE FOR
CLINICAL RESEARCH TRAINING

Course Prospectus 2023



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It is with great excitement that our new Course Prospectus for 2023 was released! With this new design and approach you will be able to navigate easily through our content and discover the services we offer for your personal growth and professional development.

You will find:

- Public courses: a variety of courses available to the public, including the course schedule for 2023
- Tailored courses: training solutions based on your request and needs
- More about ECCRT

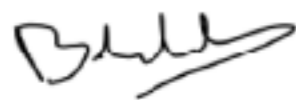
However, there is more to announce: we are transforming from a training organisation to a development organisation, with in addition to training, two additional components:

1. Increasing behavioral change after a learning intervention by using our unique methodology to stimulate self directed learning, which is key in the volatile world we live in. Read more about this on page 23.
2. Offering mentorship to ensure learnings are applied on the job with the aim of achieving clinical development excellence. Read more on page 24.

Our team is excited about this and are ready to tell you more about what we have to offer.

I wish you a pleasant reading.

Benedikt Van Nieuwenhove
Managing Director



Facts & figures

90+
PUBLIC
COURSES

30+
ONLINE
COURSES

16,000+
TRAINEES

70+
TRAINERS

in 2021

1,200+
TRAINEES

70
TAILORED
COURSES

in 2021

94%

of our TRAINEES
recommend our
courses

Our solutions

Our ultimate goal is to train talent by providing implementable knowledge in the day-to-day **activities of our learners.**

1 Public Courses

Individual & Group Solutions

We offer a range of face-to-face or live webinar training sessions scheduled throughout the year as well as courses in eLearning and microLearning formats.

Choose the training according to your needs and your preferred location and format.

2 Tailored Solutions

Business Solutions

Our Tailored Solutions are personalised training services that are provided according to your specific needs.

These solutions cover a range of areas, such as tailored courses, coaching, consulting and many other solutions, with the aim of meeting all of your training needs.

Courses categories

Hard skills

TECHNICAL TRAININGS
you can find :

- Clinical Operations
- Regulatory Courses
- Quality Assurance
- Clinical Research

Soft skills

COMMUNICATION

Training sessions for
anyone trying to improve
their communication skills.

MANAGEMENT

Training sessions for
anyone managing a
clinical trial and a team.

LEADERSHIP

Training sessions for
anyone aspiring to
become a leader.



Public courses

For individuals
and groups

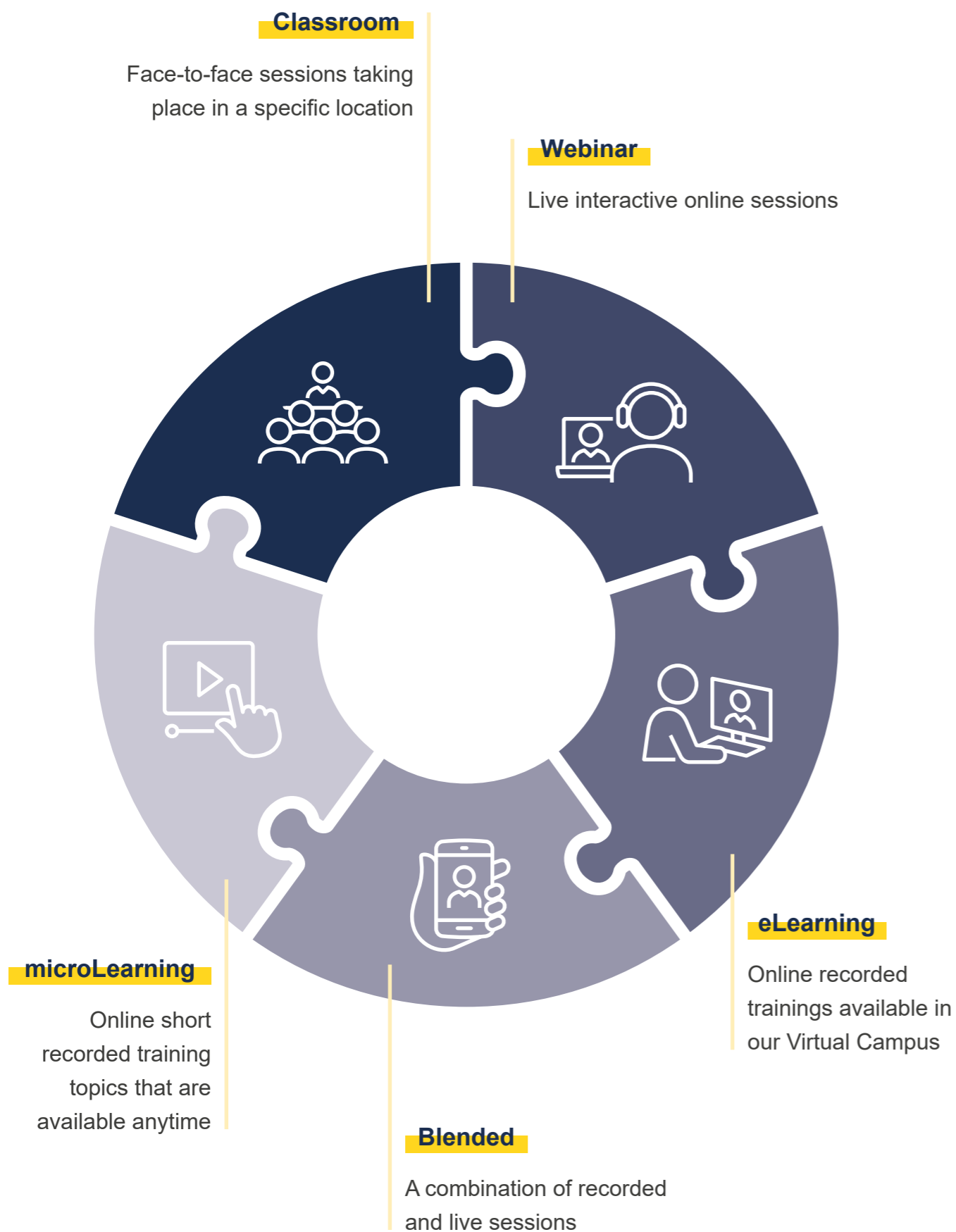
Our scheduled training programmes cover topics related to clinical research, quality assurance, regulatory affairs, as well as soft skills to make you excel in your job.

All these programmes are globally recognised and offer participants a unique learning journey through our Virtual Campus, with related course materials and official certificate.

Our courses can be accessed as classroom, live webinars and via eLearning and microLearning sessions. Some trainings are sometimes blended, i.e. carried out combining different formats.

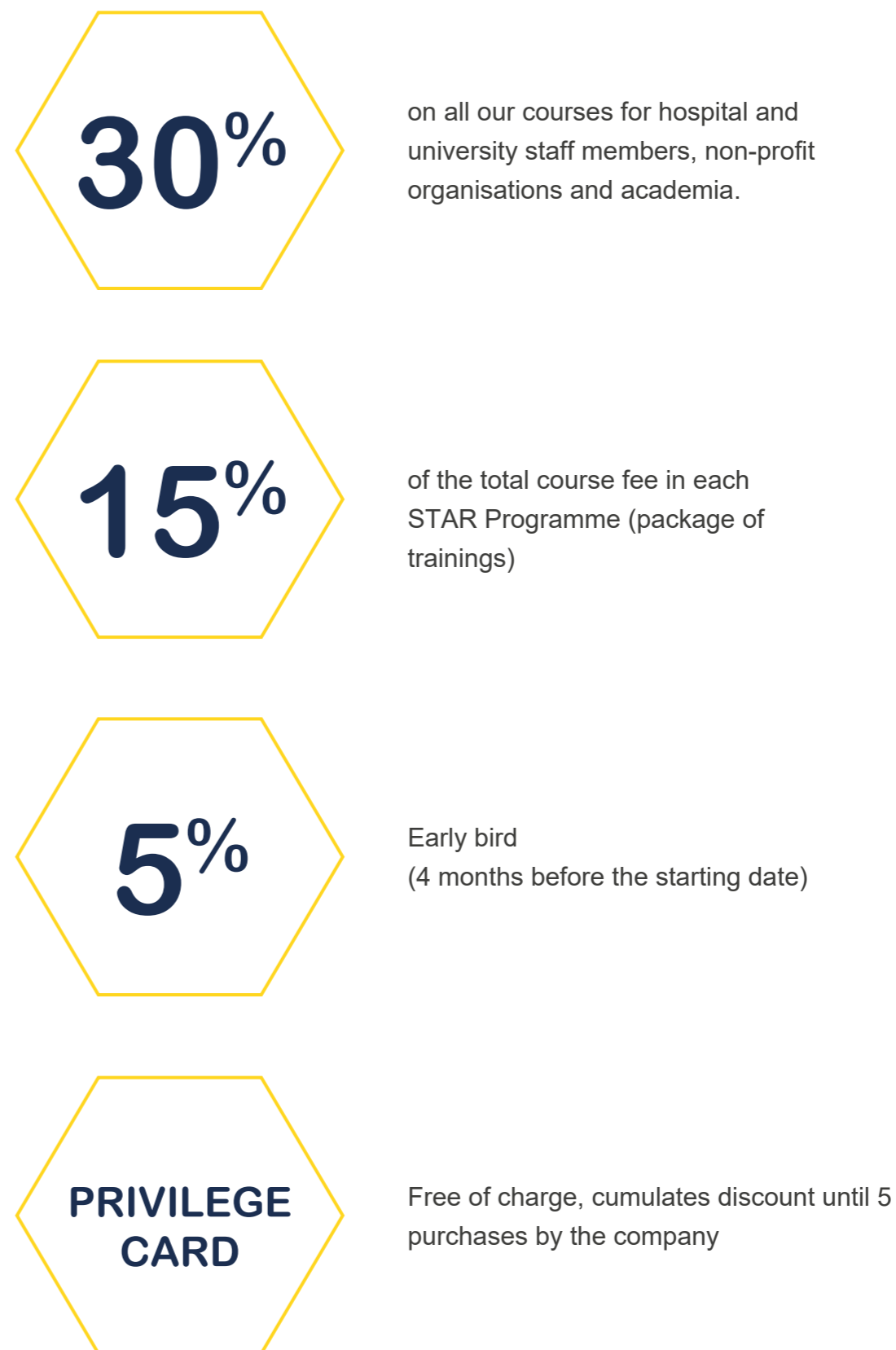
Check our course schedule to discover our training sessions planned throughout the year.

Course formats



Discount eligibility

NOTE: Discounts are not cumulative



microLearning

microLearning is a term used for small online learning units and short-term educational activities.

microLearning is a brief, to-the-point and covering practical topics such as CAPA, Investigator Site File, drug accountability log and many others, that will help you improve your productivity, avoid double work and bad time management.

At any time and anywhere, you can quickly access our short online educational videos that will provide you with the knowledge to accomplish your daily tasks at work.

microLearning categories :

- COMMUNICATION
- CLINICAL OPERATIONS
- DATA
- ETHICS
- GCP
- HOT TOPICS
- LEADERSHIP
- REGULATORY

Some of our microLearnings:

- Serious Breaches in Clinical Trials
- How to write an effective CAPA?
- eTMF Reflections
- Linking Essential Docs with GCP
- Difference between ISO 14155 and GCP
- Informed Consent the Right Way
- What to look for when not doing 100% SDV
- How To Enhance Oversight of the TMF?
- What to verify in an Investigator Site File?
- Optimising Site Selection
- Difference between Protocol Violation and Deviation
- Creating Worksheets for Clinical Source Data
- Using eSource Data in clinical investigations
- Informed Consent in a Nutshell
- How to prove oversight by the investigator?
- ICH-GCP E6 (R2) Oversight
- What is drug accountability verification?
- Applying Ethical Principles in Clinical Research
- How feasibility can make or break your study?
- Differences between TMF and ISF
- Using stakeholder's analysis to help with your project communication plan
- ALCOA - What does it stand for?
- A Systematic Approach to Safety Monitoring during Clinical Trials
- COVID-19 and ongoing Clinical Trials
- Lessons Learned
- ISO GCP Version 3! What's new?
- Project Kick-Off Strategies
- Using Social Media for Recruitment
- Resolving issues at work
- Impact and Influence
- How to deal with incomplete data?
- Vendor Oversight in Practice
- EU CTR 536



Star Programmes

STAR Programmes are a mix of both technical and soft skills training sessions, assuring that you will get an all-round curriculum, providing you with all competencies required for a specific function or role.

There are two more advantages to the STAR Programmes:

- 1 We offer you the flexibility to follow the individual courses: when you register for a STAR Programme. You can freely choose the dates of the courses throughout the year, to allow minimal disturbance of your daily activities and to optimise your travel schedules.
- 2 Secondly, the costs of the STAR Programme is 15% lower than the total price of the individual course's fees.

JUNIOR LEVEL

CLINICAL PROJECT MANAGEMENT

The perfect combination of training sessions to start or to advance in your career to become a clinical PM.

CLINICAL REGULATORY

Get an oversight of clinical study regulations with medicinal products, including the latest updates.

JUNIOR AUDITORS

The essential trainings you need to start a career as an auditor in a GCP environment.

JUNIOR CLINICAL RESEARCHER

A full week training combined with a 1-year internship to find a job in clinical research.

MEDICAL DEVICES

Learn the different products, legislations, concepts and strategies for medical devices.

REGULATORY AFFAIRS

A full week training with a 1-year internship to help you have the expertise required to start a career.

ADVANCED LEVEL

ADVANCED AUDITORS

Intended for experienced auditors who need the skills and expertise to advanced in their career.

ADVANCED CLINICAL RESEARCHER

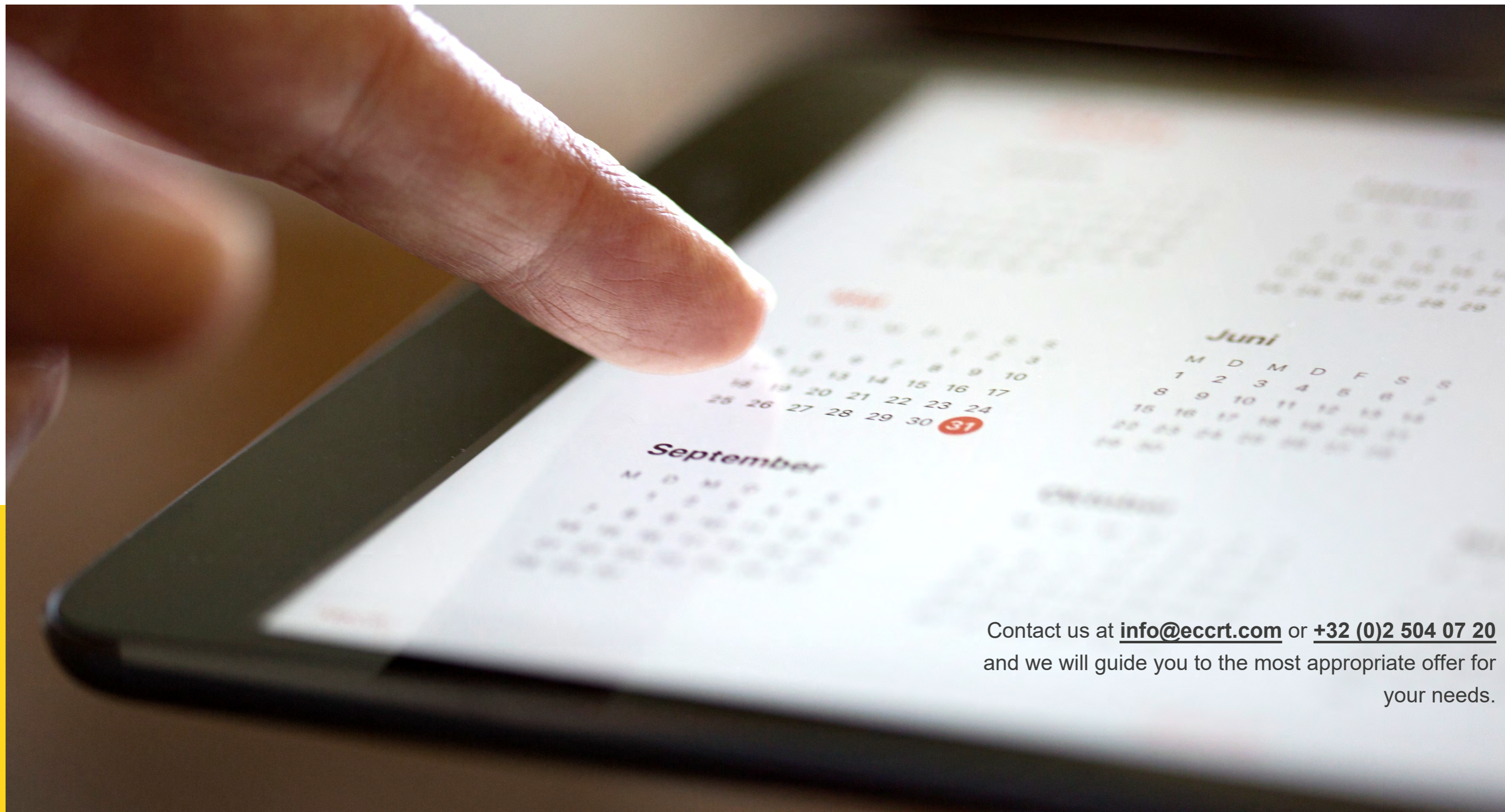
Especially for CRAs who are eager to further develop their professional soft and technical skills.

ADVANCED PROJECT MANAGEMENT

Targeted for Project Managers who wish to advance in their professional career.

Courses Schedule






Many training topics are available upon request, even if these aren't scheduled. You can either request to join our waiting list for the next session or ask for a tailored project for your organisation.






Contact us at info@ec crt.com or **+32 (0)2 504 07 20** and we will guide you to the most appropriate offer for your needs.

Course Schedule 2023

Clinical Operations courses

COURSE NAME	FORMAT	LOCATION	JAN	FEB	MAR	APR	MAY	JUN	JUL	AUG	SEP	OCT	NOV	DEC
Advanced Clinical Project Management ⁸ 	Classroom	Brussels			23, 24							16, 17		
	Webinar	Online					22, 24, 26, 30, 01, 02							
Budgeting Clinical Studies 	Classroom	Brussels						22						
Clinical Project Management ⁹	Webinar	Online			8, 10, 15, 17									
	eLearning + Classroom	Online + Brussels											7	
		Online + Zurich					12							
Clinical Research Training for Junior Clinical Research Associates ²	Webinar	Online					02, 04, 09, 11							
	Classroom	Brussels							06, 07		27, 28			
		Leiden												14, 15
Clinical Research Training for Senior Clinical Research Associates ¹	Webinar	Online				24, 26								
	eLearning + Classroom	Online + Brussels										9		
Computer System Validation for Clinical Operations	Webinar	Online			20, 22									
	Classroom	Brussels										23, 24		
CRO Management and Oversight ⁸	Webinar	Online						06, 08, 13, 15						
	Classroom	Brussels										18, 19		
Decentralised Clinical Trials - From Theory to Practice 	Classroom	Brussels										17, 18		
Designing clinical research protocols for a better outcome	eLearning + Classroom	Online + Brussels			30									
Getting the Right Level of Sponsor Oversight 	Webinar	Online						19						
	Classroom	Brussels										18		
Importance of the Involvement of Clinical Operations in Clinical Study Protocol Review ⁸	Webinar	Online						19				20		
Keeping oversight of Data Management for Clinical Project Managers 	Webinar	Online										23		
Risk-Based Monitoring ²	eLearning + Classroom	Online + Brussels										11		
	Webinar	Online					03, 05							
Risk Management in Clinical Research ¹	eLearning + Classroom	Online + Brussels											8	
	Webinar	Online			15, 17									

COURSE NAME	FORMAT	LOCATION	JAN	FEB	MAR	APR	MAY	JUN	JUL	AUG	SEP	OCT	NOV	DEC
Introduction to Clinical Research ²	eLearning	Online												
Introduction to Clinical Research with Medical Devices ³	eLearning	Online												
Managing the Trial Master File and basics of Clinical Trial Systems 	Classroom	Brussels							7					
People Management	Classroom	Brussels											9	
	Webinar	Online			29, 31									
Running Medical Device Studies ³	Classroom	Brussels						27, 28					21, 22	
Sponsor Co-Monitoring ¹	Webinar	Online					08, 10							
	Classroom	Brussels										10		
Advanced Clinical Researcher STAR Programme ¹	Classroom	Brussels										9, 10, 11		
	Webinar	Online				24, 26	03, 05, 08, 10							
Advanced Project Management STAR Programme ⁸	Classroom	Brussels										16-20		
	Webinar	Online					22, 24, 26, 30	01, 02, 06, 08, 13, 15, 19						
Clinical Project Management STAR Programme ⁹	Classroom	Brussels											6-9	
	Webinar	Online			8,9,15, 17, 22, 24, 29, 31									
Clinical Trial Assistant STAR Programme 	Classroom	Brussels							6					
Junior Clinical Researcher STAR Programme ²	Classroom	Brussels							5-7		26-29			
		Leiden												12-14
Medical Devices STAR Programme ³	Classroom	Brussels						26-28					20-22	
The following courses aren't scheduled but can be requested as a tailored course (choose topic, location, number of participants and format)														
Budgeting Clinical Trials	Upon Request													
Monitoring and Protocol Deviation	Upon Request													
GMP Essentials for Clinical Operations Staff	Upon Request													
Remote Monitoring and Opportunities during COVID-19 and Beyond	Upon Request													
Introduction to Pediatric Clinical Development 	Upon Request													
The ECG in Clinical Research	Upon Request													
Transfer of Sponsorship of ongoing trials	Upon Request													
Safeguarding Data Integrity in Highly Regulated Environments	Upon Request													
What's new with ISO 14155:2020 GCP	Upon Request													
Introduction to Statistics	Upon Request													

Course Schedule 2023



Clinical Research courses

COURSE NAME	FORMAT	LOCATION	JAN	FEB	MAR	APR	MAY	JUN	JUL	AUG	SEP	OCT	NOV	DEC
Bridging preclinical and clinical development	Classroom	Brussels											27, 28	
	Webinar	Online					02,04 09,11							
Data Protection in Clinical Research and GDPR in action	Classroom	Brussels						20						
Drug Development Cycle	eLearning	Online												
Introduction to Oncology for Clinical Researchers	Classroom	Brussels										26, 27		
Legal Basics for Clinical Study Contracts ⁹	Classroom	Brussels											6	
	Webinar	Online			22, 24									
Orienting your Career in Clinical Research ²	Classroom	Brussels							5		26			
	Classroom	Leiden												12
How to prepare your vaccine candidate for clinic	Classroom	Brussels			28									
Keeping oversight of safety for Clinical Project Managers	Webinar	Online						27				24		
Pharmacovigilance System Compliance - Medical Product Life Cycle	Classroom	Brussels			20, 21									
The following courses aren't scheduled but can be requested as a tailored course (choose topic, location, number of participants and format)														
Clinical Development of a Vaccine	Upon request													
Data Management Essentials for Clinical Project Managers	Upon request													
Implementing GDPR in your organisation	Upon request													
ISO14155 Training ³	Upon request													
Critical Literature Review	Upon request													
Data Management Essentials for Clinical Project Managers	Upon request													
Effective Medical Writing & Data Presentation	Upon request													
Introduction to Statistics	Upon request													
Laboratory Testing in Clinical Research	Upon request													
Learning Journey on Drug Development	Upon request													
Recist Training	Upon request													
Writing Procedural Documents	Upon request													

COURSE NAME	FORMAT	LOCATION	JAN	FEB	MAR	APR	MAY	JUN	JUL	AUG	SEP	OCT	NOV	DEC
A Risk-Based Approach to Clinical Audits	Classroom	Brussels						21						
Audit and System Inspection Readiness - How to be prepared ^{1 5 8}	Classroom	Brussels						14						
Auditing Clinical Development Documents ⁵	Webinar	Online						15						
Clinical Service Provider Audits ⁴	Classroom	Brussels						22						
Communication and Appreciative Auditing ⁵	Webinar	Online						16						
Introduction to System Audits for Clinical Auditors ⁴	Classroom	Brussels						19						
Introduction to Regulatory Affairs	Webinar	Online									14			
Introductory Course on Auditing Investigator Sites ⁵	Classroom	Brussels						12, 13						
Writing Audit Reports ⁴	Classroom	Brussels						20						
Advanced Auditors STAR Programme⁴	Classroom	Brussels						19-22						
Junior Auditors STAR Programme⁵	Classroom	Brussels						12, 13, 14						
	Webinar	Online						15, 16						
The following courses aren't scheduled but can be requested as a tailored course (choose topic, location, number of participants and format)														
Remote Auditing	Upon request													

Course Schedule 2023

Regulatory courses

COURSE NAME	FORMAT	LOCATION	JAN	FEB	MAR	APR	MAY	JUN	JUL	AUG	SEP	OCT	NOV	DEC
Basics on Regulatory Requirements in Clinical Research ²	eLearning	Online												
Clinical Research Training for Clinical Trial Assistants (CTAs)	Classroom	Brussels											13	
Foundational ICH-Good Clinical Practice (GCP) E6 (R2) Training ²	eLearning	Online												
GCP Essentials in 90 Minutes	eLearning	Online												
Good Manufacturing Practice (GMP) in relation to GCP	eLearning	Online												
ICH-GCP E6 (R2) Refresher	eLearning	Online												
ICH-GCP E6 (R2) Refresher + Complementary ICH-GCP Refresher for Biometrics Staff	eLearning	Online												
ICH-GCP E6 (R2) Refresher + Complementary ICH-GCP Refresher for Clinical Operations Staff	eLearning	Online												
ICH-Good Clinical Practice (GCP) E6 (R2) for Investigators	eLearning	Online												
Implementation of MDR 2017/745 in Belgium	eLearning	Online												
Impact of the EU CTR 536/2014 on your organisation 	Webinar	Online				27								
Introduction to Regulatory Affairs 	Webinar	Online									14			
Clinical Investigations: Implementation of MDR 2017/745 in Belgium	Webinar	Online				20								
How to cope with the development of drug-device combinations?	eLearning	Online												
ICH-GCP E6 (R2) Refresher + Complementary ICH-GCP Refresher for Regulatory Staff ⁵	eLearning	Online												
Local Clinical Trial Legislation in the USA ⁵	eLearning	Online												
Medical Device Regulations ³	Classroom	Brussels						26					20	
Orienting your Career in Regulatory Affairs	Classroom	Brussels						22						
The Belgian Clinical Trials Law of 2017: A Clear View on Rules	Webinar	Online			27							12		
The European Clinical Trial Directive for Medicinal Products ⁶	eLearning	Online												
The European Clinical Trial Regulation (CTR) 536/2014 ⁶	Webinar	Online						20					16	
Understanding and complying with the EU CTR	Webinar	Online					23, 25							12, 14
	Classroom	Brussels		28								5		
Clinical Regulatory STAR Programme ⁶	Webinar	Online						20					16	
Regulatory Affairs STAR Programme ⁷	Classroom	Brussels									18-22			
The following courses aren't scheduled but can be requested as a tailored course (choose topic, location, number of participants and format)														
Ready for the IVDR? Regulatory impact and milestones for CE marking	Upon request													
Clinical Investigations: Implementation of MDR 2017/745 in Belgium	Upon request													
European Legislation for Clinical Research – Implementation in Belgium	Upon request													
How to cope with the development of drug-device combinations?	Upon request													
Good Clinical Laboratory Practice	Upon request													
Good Manufacturing Practice (GMP)	Upon request													
Good Distribution Practice (GDP)	Upon request													

*Courses followed by numbers in upper script are the training sessions included in a specific STAR Programme.

Course Schedule 2023

Soft Skills courses

COURSE NAME	FORMAT	LOCATION	JAN	FEB	MAR	APR	MAY	JUN	JUL	AUG	SEP	OCT	NOV	DEC
Communication Skills ²	Classroom	Brussels									29			
Line Management Essentials	Classroom	Brussels										26, 27		
People Management ³	Webinar	Online			29, 31									
	Classroom	Brussels											9	
Career Launch Coaching (flexible schedule)	Webinar	Online	Upon your agenda											
Clinical Career Coaching (flexible schedule)	Webinar	Online	Upon your agenda											
The following courses aren't scheduled but can be requested as a tailored course (choose topic, location, number of participants and format)														
Change Management for Clinical Research	Upon request													
Communicating with EU Regulators/Health Authorities	Upon request													
Enhancing your Communication and Presentation Skills in the changing Clinical Trial world	Upon request													
Influencing skills	Upon request													
Leading in a Solution Focused Way	Upon request													
Stress Prevention at Work	Upon request													
Time Management	Upon request													
Microsoft Project Basics for Clinical Project Managers	Upon request													
Intercultural Communication Skills	Upon request													
Remote Team Management : How to successfully lead	Upon request													
Train the Trainer	Upon request													
Female Leadership in Clinical Research	Upon request													

Tailored Solutions

For organisations



Tailored Solutions

We offer a range of possibilities to develop training sessions according to your expectations and needs. You can opt for:

- Tailored courses
- Coaching
- Consulting
- Competency Framework
- Team To Go
- Behavioural Change
- Clinical Development Excellence



Tailored Courses

Request any training about a clinical research related topic for your organisation. You can decide:

- **Topic**
what points to focus on that subject
- **Training format**
face-to-face or live webinar and via eLearning sessions. Some trainings are sometimes blended, i.e. carried out combining different formats
- **Location of the training**
- **Number of participants**



Competency Framework

Competency Framework is a standard mapping of competencies to roles, used to help organisations assess and manage individual and collective work performance.

Based on the developments of the Joint Task Force for Clinical Trial Competency and on our vast expertise in Clinical Research, ECCRT has taken the Competency Framework concept a step further to fit your specific needs.



The Competency domain includes:

- **Scientific Concept and Research Design**
- **Ethical Participant Safety Consideration**
- **Clinical Studies Operations (GCPs or ISO 14155)**
- **Study and Site Management**
- **Leadership and Professionalism**



Team To Go



Core team meeting

- Define the team vision and means
- Define goals and objective
- Define KPI

Teaming

- Team building and workshops to enhance team dynamics & performance

Guiding & Coaching

- Ongoing training via an online forum and Q&A sessions
- Coaching individuals, including performance management and competency assessment

The Team To Go programme focuses on your team and on the communication between each stakeholder in order to increase efficiency in the day-to-day activities of a team.

Team To Go (TTG) is not only about training a team on specific skills, but it is also about bonding the team for success. Not designed to be a one-time effort, it will provide guidance and coaching to your team throughout the project in order to have a successful outcome.

The main 3 components are:

Link: <https://ecrt.com/team-to-go>

Coaching

At ECCRT, we aim at guiding you and your team to succeed and to brilliantly face new challenges. Therefore, we have put different coaching solutions in place that will be beneficial for you: career, personal and team coaching. Together with our expert coaches you will achieve your goals.

You can choose between:

- Coaching on how to **start your career** in Clinical Research¹
- Coaching on how to **advance in your career** in Clinical Research²

Consulting

ECCRT is your ideal partner to provide consultancy about any matter related to the acquisition of competencies. Whether this is about how to set up a training department, how to develop your team's technical skills or how to build a competency framework, we are there to help you be successful.³



¹ Link: https://eccrt.com/course_display/career-launch-coaching/

² Link: https://eccrt.com/course_display/clinical-career-coaching/

³ Link: <https://eccrt.com/consulting>

Behavioural Change



BEHAVIOURAL DIMENSION

Did you find difficulties in following-up with a training? Do you struggle with changing the performance and activities of your staff?

ECCRT provides training and development programmes that combine conventional training tools and approaches with innovative interventions rooted in evidence from behavioural sciences and tailored to the client's circumstances and requirements.

What does it imply?

Behavioural dimension to Learning and Development, through galvanising the entrepreneurial potential of employees

EMPIRICAL RESULTS

Creation and monitoring of data-driven and evidence-informed approach based on the principles of design, behavioural and educational sciences principles

INNOVATIVE LEARNING

Areas such as stimulating or revitalising innovation, ensuring compliance, strengthening entrepreneurial and entrepreneurial culture, designing robust Learning and Development interventions

Clinical Development Excellence

Journey to Excellence

Reducing risks

We commit to lowering the risk of your clinical developing failing because, as clinical research experts, we help you tackle the challenges you face such as delayed timelines, lack of quality, high costs etc.



Team development

Our experienced mentors accompany you throughout the clinical development process and assure the streamlined involvement and development of your teams.

Is your team lacking engagement or facing inefficiencies? Have you ever noticed insufficient conformity or quality? Are you often confronted with uncontrolled financial losses due to the risks faced during clinical development?

Our mentors will accompany your teams throughout the journey of the clinical research process with a 360° development approach.

More than just expertise, it is all about mentoring

TECHNICAL EXPERTISE

Thanks to our strong network of experts, we can clearly identify gaps and propose high-quality training, to build the foundation of required knowledge.

ON-THE-JOB TRAINING

Our experienced mentors will accompany your staff in their on-the-job challenges in order to consolidate the previously acquired concepts and deepen their operational background.

COMPETENCIES DEVELOPMENT

Ultimately, your staff will develop their competencies, become more effective and increase the value of your organisation.



About ECCRT

ECCRT Mission
Lab
Trainers Network
Testimonials
FAQ

ECCRT Mission

We are a professional Clinical Research Training Organisation for the **pharma, biotech & medical device industries** as well as for CROs, investigational and academic groups.

Our mission is to facilitate Clinical Research professionals to **excel in their job** for the benefit of patients.

To achieve this, we have **70+ seasoned experts on our team** covering a wealth of know-how which they enthusiastically share with our course participants in the most interactive, engaging and innovative way.



WHY ECCRT?

EFFECTIVE LEARNING

Our goal is to provide a **360 degree approach** combining training, coaching, and consulting, benefiting individuals and organisations!



HUMAN APPROACH

We want you to be trained by **real and experienced trainers** who are keen to share and personalise their trainings according to your needs!



INNOVATION

With our Virtual Campus, you can **learn, share and exchange knowledge** as you have never experienced before!



ECCRT Lab

What is the ECCRT Lab?

The ECCRT Lab was created with the intention of understanding your needs, establishing a way to better interact and sharing ideas for future training development.

The lab is open to anyone looking to engage with us and to help us shape the future of clinical research for the benefit of patients.

Shape the future of your training now. With this platform, you can share your development needs, see what ECCRT is developing, and join exploratory sessions!

What to expect?

- New courses
- Surveys
- Course catalogue
- Discovery sessions

Our trainers



Aimad Torqui



Amer Alghabban



An Vijverman



Angeles Escarti-Nebot



Anja Schiel



Anne Miermont



Anya Nijenhuis



Ann Lampo



Anne Miermont



Anne Lenaers



Barbara Gastl



Bart Derre



Benedikt Van Nieuwenhove



Begonya Nafria Escalera



Bodo Lutz



Dimitrios Athanasiou



Dominique Monferrer



Elisabeth Reus



Eline Detobel



Eric Rozet



Essam Ghanem



Greet Musch



Hedwig Beernaert



Hugues Bogaerts



Jean Van Rampelbergh



Johan Bosmans



Johannes Kreuzer



Jolanda Schavemaker



Joris Vandeputte



Karen Gabriels



Karel Allegaert



Kevin Punie

Our trainers



Koen Norga



Kristof Vercruysse



Kroen Nauwelaerts



Liesbeth Lemmens



Lourens Bloem



Ludivine Petit



Ludwig Everaert



Marc Devisch



Marc E. Brooks



Marisa Giro



Marleen Verbeeck



Mieke Tempels



Morgane Franck



Mustafa A. Zaman



Nancy Cottigny



Nathalie Niclaus



Nelle Stocquart



**Olivier Van
Obberghen**



Paula Hemdal



Peter Musschoot



Pierre Van Damme



Pieter Wyckmans



Pieter Vancaeneghem



Pieter Neels



Robert Edwards



Roman Bobrovsky



Saskia De Haes



Sandrine Tinton



Sini Eskola



Steven Thys



Thomas Ockier



Ruth Beckers

Our trainers



Tim De Schutter



Tom Van Paepegem



Vincent Baeyens



Virginie Hamtiaux



Wim De Meester



Zuzanna Kwade

ARE YOU INTERESTED IN BECOMING A TRAINER?

Join us

ECCRT has a broad network of trainers, coaches and subject matter experts, always providing you with the best and most up-to-date insights and hands-on learning. To keep providing valuable content to students, we are constantly looking for new trainers to share their expertise.

Contact us at info@eccrt.com
or **+32 (0)2 504 07 20** and
we'll work together on creating a great learning experience.

Testimonials



This week has been intense and very rich in learnings. The richness of this training comes by large from the speakers/teachers who bring their real-world experiences of the field and their vision of the regulatory history and future changes in Europe...

- **Juliette Moyerso** (Regulatory Affairs STAR Programme)



It was very helpful to get an overview on all aspects to be considered in regard of pre-clinical to pre-clinical phase, especially the regulations referred to. So it's easier to read through the one or the other part later on when required.

- **Michaela Schaden** (Bridging Preclinical and Clinical Development)

Thanks to the Jr Clinical Researcher STAR Programme I understood better who can become a CRA and where to go. Through this knowledge and the interaction I had with the trainer, I know how to start and build my career in clinical research.

- **Braulio Lima** (Junior Clinical Researcher STAR Programme)



The training was very well explained by the trainer, and the work materials were very interesting and retrievable for future questions. I got a more precise idea of the PM's role in clinical project management.

- **Frederique Jacquinot** (Clinical Project Management)



I decided to take the ECCRT career coaching. This helped me get a better overview of the roles and functions in the area of clinical development. Together with my coach – Virginie Hamtiaux – we identified what my own strengths are and how to link them to a possible career in the pharma industry.

- **Kristien Van Belle** (Career Coaching)



Insightful lectures, friendly environment and interactive learning experience. I was very welcomed by the ECCRT team who provided a pleasant and well-located place to do the training. Lectures were given by highly experienced professionals who gave a clear understanding of the drug life cycle and the required regulation...

- **Joana Reis** (Regulatory Affairs STAR Programme)



I found the course extremely helpful. The slides and content were very well structured and presented and Kirsten simply was amazing as a trainer and presenter. I also liked Sanofi's presentation sharing experience made working in CTIS.

- **Christian Riegel** (Understanding and complying with the EU CTR)



The deep expertise of the presenters, while still being accessible, speaking in a comprehensive matter. No answer was made up. Great answers, exactly on the questions. The whole program «tells a story». Not just a bunch of slides put side by side. The best training session I had in a long, long time.

- **Federico Melo Ferrer** (Introduction to Oncology for Clinical Researchers)

Liesbeth Lemmens did such a great job in training us online and interactively that I completely got the classroom feeling! Well done Liesbeth and thank you very much! It was a great experience and so so so fruitful and helpful!

Great teacher, great teammates!

- **Regula Schneider** (Clinical Research Training for Junior CRAs)



The Programme was the perfect opportunity for me to gain practical experience in this field. The internships in 3 different organisations gave me a better view of the different job functions and possibilities. I'm ready to dive into the world of clinical research...

- **Veronique Shiwa** (Junior Clinical Researcher STAR Programme)



Questions & Answers

1. What is the difference between our Foundational GCP training and the GCP Refresher training?

The **Foundational GCP training** is a course on all chapters of the ICH-GCP guideline E6. We focus on the ethical aspects of performing clinical trials outlining the responsibilities of all stakeholders. Through extensive theory and interactive moments, including real life situations, we will get you acquainted with the ins and outs of Good Clinical Practice. You can either choose to do it **online** or as a **face-to-face session**.

The **GCP Refresher training** is designed for people already working in a GCP environment and just needing an update on this topic. During this course, we immerse you with examples of day-to-day situations and guide to successful solutions of these problems.

Please look at our catalogue to select the best option for you.

If needed, you can always request a training for your company or team, as we will tailor the training according to your needs.

[Contact us for more information.](#)

2. How do I access the online grading test for my course?

1. Log into eccrt.com with your credentials
2. Click 'Virtual Campus' in the top left hand corner of the website
3. Click 'Enter the Virtual Campus'
4. Then you should see your course on your Dashboard
5. Click on your course and then go to the tab "Take the final test".

3. Do we receive a certificate at the end of the course?

Everyone who participates in our courses receives a **certificate of attendance**. We also offer a **Certificate of Achievement** for a growing selection of our courses.

The Certificate of achievement can only be given to participants who completed their test with a score of at least 70%. Participants are allowed to consult their course notes and the extensive training material provided.

In the case of an online grading test, you will be offered to download your digital certificate directly from the ECCRT Virtual Campus once you've completed your test.

[All certificates are globally recognised.](#)

4. What is ECCRT's cancellation policy?

If you need to cancel your registration, the following cancellation terms are applicable:

- Cancellation by the ECCRT: due to unforeseen circumstances, it is possible that the training programme may change and the ECCRT reserves the right to alter the venue or to cancel the event.
ECCRT reserves the right to reschedule or cancel classes up to two weeks prior to the scheduled date. Registered participants will be informed of any such changes.
- Cancellation by the registrant:
 - More than 30 calendar days prior to the course: no cancellation fee will be charged.
 - Between 21 – 30 calendar days prior to the course: 50% of the fee will be charged.
 - Less than 21 days prior to the course or if no notification is received: the registrant will be liable to pay the full course fee.
 - An administration fee of 50 EUR will be charged for each cancellation.
- Alternative delegates may replace a registrant; however all cancellations/replacements must be received in writing.

[Read more about our terms and conditions.](#)



EUROPEAN CENTRE FOR
CLINICAL RESEARCH TRAINING

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