

CLINICAL RESEARCH

WHAT'S IT ALL ABOUT?

2022
EDITION

...a focused training for busy investigators and study teams.

WHEN:

21 & 22 November 2022 | Online

COURSE DIRECTORS

Inês Zimbarra Cabrita, PhD
Francisca Patuleia Figueiras, PhD

SCIENTIFIC COMMITTEE

Fausto J. Pinto, MD, PhD, FESC, FACC, FSCAI, FASE
Dulce Brito, MD, PhD, FESC
Joaquim Ferreira, MD, PhD
Catarina Sousa, MD, PhD
Cristina Valente, PharmD

TARGET AUDIENCE:

Investigators, pharmacists, study nurses and study coordinators with zero to 5 years of experience in clinical research and anyone who would like to learn the fundamentals of Good Clinical Practice and extend their knowledge in clinical research methodologies and procedures.

DESCRIPTION:

Running clinical studies is a complex task that requires several skills. Skills ranging from Good Clinical Practice over all applicable regulations up to operational aspects on how to carry out clinical studies. Having a high trained and specialized study team conducting clinical research is the main key to achieving success in recruitment objectives and high standards of quality and performance.

This interactive 2-day course will provide you a comprehensive knowledge on the practical aspects of clinical studies, essential to reach the highest quality of data whilst ensuring the study participants' safety and well-being and that your professional knowledge is optimized.

ACCREDITATIONS



ORGANIZED BY

CETERA, a Portuguese Academic CRO, an Autonomous department of the Association for Research and Development of the Faculty of Medicine, based at the Lisbon Academic Medical Centre (CAML).

PROGRAM | Main Topics

DAY 1

9am – 5pm

- Session 1. Types of Studies and Research Design
- Session 2. Regulatory Aspects
- Session 3. Clinical Study: the first contact with investigators and study team
- Session 4. Patient-centered Research
- Session 5. Interactive Workshop

DAY 2

9am – 5pm

- Session 6. The Physician as Clinician and Principal Investigator
- Session 7. Clinical Study Ongoing Activities & Stakeholders
- Session 8. Audits and Inspections
- Session 9. Pharmacovigilance
- Session 10. Interactive Workshop



IN COLLABORATION WITH



SCIENTIFIC SPONSORS



STAY CONNECTED WITH US!

AIDFM, Av. Prof. Egas Moniz
Piso 01, 1649-028 Lisboa, Portugal
training.cetera@medicina.ulisboa.pt
Telf.: +351 21 793 09 20
www.aidfm-cetera.com

