

CLINICAL RESEARCH:

WHAT'S IT ALL ABOUT?

2020

EDITION

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

WHEN

Next Editions

15 & 16 October, 2020

COURSE DIRECTORS:
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SCIENTIFIC COMMITTEE:
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TARGET AUDIENCE:

Health Care Professionals (doctors, allied health professional, nurses, study coordinators, administrative assistants) 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.

ACCREDITATION



ORGANIZED BY
AIDFM-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. Patient-centered Research
- Session 4. Clinical Study: the first contact with investigators and study team
- 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:



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