

CERTIFIED CLINICAL RESEARCH PROFESSIONAL TRAINING COURSE



This program is designed to provide training in ICH-GCP guidelines, clinical trials monitoring, investigative site coordination, understanding local/ international regulations, and risk management in clinical trials.



Module:

12 Online/Onsite sessions covering a comprehensive curriculum of clinical research.

Eligibility:

M.B.B.S/ B.D.S, registered nurses, medical students, M.Phil./ Ph.D. B.Pharm. / Pharm-D/ M.Pharm. B.Sc. (biological sciences), M.Sc. (biological sciences)

Assessment:

Total 10 assessments (40% weightage)
One final exam (60% weightage)

Timing:

Every Sunday 10:00 AM to 01:00 PM.

Batch-wise Planned Dates:

Batch 01	Batch 02	Batch 03	Batch 04
Februrary	May	August	November
2024	2024	2024	2024

REGISTER NOW

VISIT OUR WEBSITE:
www.mrcro.com/ccrp/
+92-21-37224982

Location for Onsite Training:

Plot # B-10, Block-16, W.C.H.S, K.D.A Scheme No.24, Gulshan-e-Iqbal, Karachi, Pakistan



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COURSE OUTLINE

DETAILS	MODULES		
Pharmaceutical Industry Overview, Drug Discovery & Research.	 Introduction to Pharmaceutical Industry Drug Discovery and Development The Pre-Clinical Studies Phases Of Clinical Trials 		
Clinical Studies, Design, & Clinical Trials	 Types of Clinical Research Essential Elements of Clinical Trials Clinical Trial Design 		
IRB & Regulatory Process	 Drug Approval Process IRB/IEC roles and responsibilities FDA Review Process Trial approval process (DRAP) 		
Historical events leading to Formation of ICH-GCP – E6 R2 Guidelines	 Historical Events leading to the current code of Bio-Ethics in Clinical Research Evolution of ICH-GCP ICH-GCP E6 R2 guidelines and updates in E6 R3 		
Clinical Trial Management	 Pre-study and Study Set-up Activities Site Selection & Site Initiation Visit Clinical Conduct Subjects Recruitment Retention Compliance Safety Reporting & Source Documents Verification 		
Essential Documents	 Overview of Clinical Trial Essential documents Study Protocols Informed Consent Process & Elements of ICF CRF and design of CRF Data Collection and Management in Clinical Trials EDC Tools 		
Stakeholder Management in Clinical Research/ Trial	 Sponsors and Stakeholders Clinical Trial Sites and Stakeholders CROs and Stakeholders 		
Risk Management in Clinical Trials	 Identification of risks and issues Mitigation plans and Review of status for mitigation. 		
Statistical Issues in Clinical Trials/ Clinical Research	 Study Design Research Methodology Statistical Consideration in Clinical trial Null & Alternative hypothesis Descriptive V/S Analytical Studies Prevalence Studies & Case Studies 		
SPSS Hands-on Training	 Data Handling Data Management SPSS – hands-on 		
Bio-ethics in a Clinical Trial	 Understand the difference between ethical and unethical clinical conduct in various scenarios: Informed Consent, Authorship, Therapeutic Misconception, Clinical Equipoise, Use of Placebo in Clinical Trials 		