

## CTR 115 Clinical Research Regulations

### COURSE DESCRIPTION:

Prerequisites: CTR 110

Corequisites: None

This course covers the range of national and international regulations and guidances governing the development of drugs, diagnostics, medical devices, and biologics. Topics include a review of the regulatory agencies, guidelines for regulatory application, required documentation, and protection of human subjects. Upon completion, students should be able to demonstrate a basic understanding of regulations and guidelines associated with clinical research and describe effective means of compliance. Course Hours Per Week: Class, 3. Semester Hours Credit, 3.

### LEARNING OUTCOMES:

Upon completion of this course, the student will demonstrate basic cognitive and practical knowledge and skills in each of the following areas:

1. Apply regulatory, legal, and guidance parameters within a clinical research project
2. Identify the function, composition, activities, and responsibilities of an Institutional Review Board (IRB)/Independent Ethics Committee (IEC)
3. Describe the components of the informed consent process
4. Define the contractual relationships and party obligations in a clinical research project or study
5. Identify the regulatory and legal documents associated with a clinical research project or study
6. Describe scientific misconduct and its consequences

### OUTLINE OF INSTRUCTION:

- I. Apply regulatory, legal, and guidance parameters within a clinical research project
  - A. Identify the domestic and international regulatory agencies and branches that impact on the conduct of clinical research
  - B. Describe the process of regulatory compliance within the context of clinical research, including ICH Guidelines, IND and IDE regulations, NDA, BLA, 510(k), and PMA submissions, marketing approval and post-marketing surveillance
- II. Identify the function, composition, activities, and responsibilities of an Institutional Review Board (IRB)/Independent Ethics Committee (IEC)
  - A. Describe the requirements for membership on an Institutional Review Board according to Federal regulations and ICH guidelines
  - B. List the specific criteria that must be fulfilled for at least five members of the Institutional Review Board
  - C. Describe how Institutional Review Board records are maintained and retained
  - D. Identify the standard review criteria used by an Institutional Review Board
  - E. Define "expedited review criteria"
- III. Describe the components of the informed consent process

- A. List the basic elements of an informed consent form according to Federal regulations and ICH guidelines
  - B. Evaluate a written informed consent form for compliance with the eight basic elements
  - C. Describe the methods that can be used to obtain informed consent as indicated in the regulations
  - D. Identify the circumstances when it is not necessary to obtain informed consent
- IV. Define the contractual relationships and party obligations in a clinical research project or study
- A. Describe the contractual nature of the FDA form 1572 as well as the components of this form
  - B. Define the obligations of sponsors and investigators in conducting clinical research under Federal regulations and ICH guidelines
  - C. Describe the obligations that can be delegated, to whom they can be delegated, and the appropriate methods for documenting the delegation
- V. Identify the regulations and regulatory documents associated with clinical research
- A. Differentiate between an adverse event and a serious adverse experience; describe the process for reporting adverse events and serious adverse experiences, including the use of form FDA 3500A
  - B. List the requirements/guidelines set by the FDA for facilities participating in clinical trials
  - C. Identify the essential documents required by regulations
  - D. Identify the regulations pertaining to shipping investigational product
  - E. Identify the regulations pertaining to the clinical research protocol and its amendments
- VI. Define scientific misconduct and its consequences
- A. Describe the ethical and legal consequences/sanctions that can be conferred on individuals and/or institutions found guilty of scientific misconduct

#### **REQUIRED TEXTBOOK AND MATERIAL:**

U.S. Federal Regulations available online at [eCFR :: Home](#) and International Conference on Harmonization Guidelines for Drugs, Biologics, and Medical Devices available online at [ICH Official web site : ICH. <http://www.ich.org/cache/compo/276-254-1.html>](#)

#### **SUGGESTED REFERENCES, PERIODICALS, AND VISUAL AIDS:**

IRB Information Sheets available online at [Search for FDA Guidance Documents | FDA](#)

Optional Textbook: Book CG1-IC: 2023 Comprehensive Clinical Research Desk Reference for Drug and Medical Device Trials (With New FDA Informed Consent Guidance). Available online at:

<https://www.clinicalresearchresources.com/products/book-cg1-2023-comprehensive-clinical-research-desk-reference-for-drug-and-medical-device-trials>