Responsibilities of the Research Team

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A successful clinical trial program is typically comprised of a team of individuals who are committed to conducting high-quality clinical trials. All staff involved in clinical research must adhere to the regulations and understand the guidelines that govern clinical research. All research teams are led by a Principal Investigator (PI) and have research participants. Other key staff include, but are not limited to: Nurses, Study Coordinators, and Clinical Data Managers. This module will provide an overview of key roles of the research team. The module is divided into 3 required components (videos) and 3 recommended videos. Once you have completed all required parts, please link to the evaluation after which you will be directed to a certificate of completion for your education records.

Module Objectives

At the conclusion of this module, learners will be able to:

- Describe the role and responsibilities of the Investigator as described by OHRP, FDA, and ICH GCP
- Describe the role and responsibilities of the Research Nurse.
- Describe the role and responsibilities of the Study Coordinator.
- Describe the role and responsibilities of the Clinical Data Manager.
- Describe the role and responsibilities of the pharmacist.

Part 1	Role and Responsibilities of the Principal Investigator (PI) (24½ minutes)	Part 1	Resources	Evaluation
Part 2	Responsibilities of the Clinical Trials Team (15 minutes)	Handout Part 2 Handout		
Part 3	Clinical Research Professional Organizations (7 minutes)			
		Part 3 Handout		
Reco	mmended			

- Clinical Trial: A Nursing Perspective (19 minutes)
- The Role of the Research Nurse (10 minute)