IRBMED Education & Training Program

OVERVIEW

IRBMED’s Education & Training Program was designed and developed through collaboration with members of the research community. It is our goal to make sure this program is tailored to meet the ongoing educational needs of the research community.

LEVELS

Classes will be structured to provide varying levels of information based on the needs and experience of the individual.

Level 1: Will offer *fundamental* courses designed to provide a broad understanding of medical research.

Level 2: Will offer *intermediate* courses designed to provide a more comprehensive understanding of the subject matter.

Level 3: Will offer *advanced* courses designed to provide information related to specific subject matters.

Level 4: Will offer *specialized* courses designed to address very specific and limited circumstances

SUBJECT GROUPS

Course offerings will be structured in subject groups designed to increase in depth and complexity from one level to the next.

Agreements: Information related to various inter- and intra-institutional agreements.

Consent: Information regarding regulatory practices, as well as technical information regarding constructing ICDs.

eResearch: Information for working within the electronic application submission system.

Non-Standard Applications: Information regarding when and how to utilize non-standard applications within eResearch.

Privacy and Confidentiality: Information related to both the regulatory aspects of privacy and confidentiality, as well as the technical aspects of storage and retention.

Regulations: Various regulatory requirements placed on research, with particular emphasis on FDA, OHRP, and OCR.

Reporting: Variety of topics related to reporting, in relation to institutional policies and external oversight entities.

Specialty: Miscellaneous topics, such as data coordination, deception research, and study team responsibilities.