Human Subjects Research and the IRB

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Overview:

- The IRB and Regulatory Oversight
- Protocol submission to the CWRU IRB
- SpartaIRB system
- Available Resources
Institutional Review Board (IRB)

- An independent group that is charged with reviewing research to ensure that human subjects’ rights and welfare are adequately protected.
- Consists of:
  - Scientists
  - Non-scientists
  - Community or non-affiliated representative
  - Men/Women
  - Ethnic/Cultural diversity
  - Prisoner representative
History of Research Abuses

- Nazi Experiments (WWII)
- Tuskegee Syphilis Study (1932-1972)
- Thalidomide Tragedy (1960)
- The Milgram Study (1963)
- Willowbrook Study (1963)

- Common Thread: The participant did not freely and knowingly volunteer for the research because pertinent information was withheld that, if known, would have compelled a refusal to participate in the study.
Ethical Guidance

- Nuremberg Code (1947)
- Declaration of Helsinki (1964)
- The Belmont Report (1979)
  - Respect for Persons
    - Informed Consent
    - Protection of vulnerable populations
  - Beneficence
    - Risk/benefit analysis
  - Justice
    - Equitable selection of subjects
The Belmont Report Codified
Federal policy for the Protection of Human Subjects -
“The Common Rule” (45 CFR 46)
Office of Human Research Protections (OHRP)

- Establishment of Institutional Review Boards (IRBs)
- Institutional assurance of compliance
- Informed Consent Requirements
Definitions

- **Human Subject** (OHRP)
  - Living individual
  - About whom an investigator conducting research obtains **either**:
    - Data through intervention or interaction with the individual **OR**
    - Identifiable private information

- **Research**
  - Systematic investigation (includes research development, testing and evaluation)
  - Designed to develop or contribute to generalizable knowledge
Criteria for IRB Approval

- Risks minimized
- Risk-benefit ratio reasonable
- Equitable selection of subjects
- Informed consent sought and documented
- Privacy/confidentiality of data
- Additional protections for vulnerable populations
HIPAA

- The Health Insurance Portability & Accountability Act (HIPAA) is a Federal law that helps to protect the privacy of health information.

- Protected Health Information (PHI):
  - any individually identifiable health information transmitted or maintained in a medical record paper or electronic,
  - designated data set that was created, disclosed, or used in the course of providing a health care service such as diagnosis, payment or treatment.
  - 18 Identifiers considered PHI under HIPAA (http://case.edu/research/resources/forms-policies/)

FDA

Drugs and Devices
CWRU IRB
Purpose and Responsibilities

- Protect human subjects
- Support and facilitate the ethical conduct of human subjects research
- Assure institutional compliance with regulatory agencies
- Assist investigators in complying with ethical and regulatory standards
Types of IRB Review

- **Exempt**
  - Determination made by the IRB Office
  - Not the same as Not Human Subjects Research
  - Not exempt from oversight, just from federal regulations

- **Expedited**
  - Not greater than minimal risk.
  - Review by one IRB member; does not go to a meeting

- **Full Board**
  - Anything that doesn’t fit the above
  - Must go to a meeting
Protocol should be completely thought out before submitting.

- Be specific: recruitment, consent, study procedures, data security
- Download the most current version of the protocol template
- Don’t rush the submission, review documents for typos and grammar. The information must be understandable by someone outside of your field.
- All documents must have consistent information
COVID-19 Restrictions

- All study procedures that can be done remotely, should be done remotely
- Requests can be made for in-person research activities, both on campus and at locations off campus
  - Dean must first provide approval for in-person research
  - A safety plan must be completed and submitted to the Office of the President and approved prior to submission to the IRB
- On campus research can only include participants who are already on campus
Category 2: Research that only includes interactions involving educational tests (cognitive, diagnostic, aptitude, achievement), survey procedures, interview procedures, or observation of public behavior (including visual or auditory recording) if at least one of the following criteria is met:

- (i) the information obtained is recorded by the investigator in such a manner that the identity of the human subjects cannot readily be ascertained, directly or through identifiers linked to the subjects; or
- (ii) Any disclosure of the human subjects' responses outside the research would not reasonably place the subjects at risk of criminal or civil liability or be damaging to the subjects' financial standing, employability, educational advancement, or reputation

- IRB will determine if data collected is identifiable and if the information is sensitive
Survey Procedures

- Points to Consider if information being asked/provided is “sensitive” in nature:
  - If using a platform such as REDCap or Qualtrics, will the survey be collecting IP addresses?
  - If not direct identifiers are being collected, are enough demographic questions asked so that an individual might be identified through deduction?

- Surveys that do not meet the criteria for exemption will be reviewed via an expedited review, which will require that the consent form contains all the required elements as detailed in the regulations.
Informed Consent Process

Begins with recruitment
- Advertisements must be approved
- May need letters of cooperation
Informed Consent Process

Consent Document
- Template in SpartaIRB Library
- Be sure to:
  - Delete sections that do not apply
  - Remove instructional language
Informed Consent Process

**Protocol Document**

- If a waiver of consent or a waiver of documentation of consent is being requested, appropriate justification must be provided.
Data Security

- Collection
  - Remote: recommend Qualtrics, REDCap, Zoom
  - If additional data will be collected
- Transfer/Sharing
  - Import and export of data
- Storage
  - PHI and identifiable data
- Destruction
• **Anonymous**
  - Data that cannot be identified, and never could be
  - Not the same as a file from which you removed identifiers. If you have access to identifiers at any point, it is not anonymous.

• **De-identified**
  - A dataset that had elements removed so it is no longer identifiable.
  - If you have a file that is linked to identifiers with a key you have access to, your study data *is not de-identified.*

• **Identifiable**
  - A data set that contains identifiers, can easily be re-identified, or is linked to identifiers with a key
  - Includes coded data
Expedited Review

Category 5: Research involving materials (data, documents or specimens) that have been collected or will be collected solely for nonresearch purposes (such as medical treatment or diagnosis).

- Chart Reviews
  - Must be clear on how data will be recorded (if you have the ability to go back and obtain additional information from a chart, it is not de-identified). Data collected from charts is never anonymous.
Other Protocol Considerations

HIPAA authorization
  • Health Insurance Portability and Accountability Act
  • Protected Health Information

• If research will involve obtaining data from an electronic chart, then you will either need written HIPAA authorization from the participant or approval of a waiver of HIPAA authorization from the IRB
  • Consent for treatment does not meet the regulatory requirements for consent for use of data/specimens in research
SpartaIRB System

- Individuals do not automatically have an account
- Help Center and Library
- Help/Information bubbles

spartairb.case.edu
Common Mistakes

- PI cannot be a student/resident
- Upload documents in correct sections
- Funding question must have an answer
- “Stack” Documents
What else can help your IRB process?

- If you receive request for clarifications:
  - Do not send back to the IRB until all changes are made
  - If you aren’t sure, contact the IRB administrator (use comment)
- All study members are required to:
  - Complete CREC Certification
  - Submit Conflict of Interest Disclosure
Available Resources:

- **IRB general email** cwru-irb@case.edu
- **CWRU IRB Policies** - [https://case.edu/research/faculty-staff/compliance/cwru-policies-and-procedures](https://case.edu/research/faculty-staff/compliance/cwru-policies-and-procedures)
- **Utech-** [help@cwru.edu](mailto:help@cwru.edu) https://case.edu/utech/
- **REDCap-** Sheree Hemphill, sheree.hemphill@uhhosptials.org
- **Case.box.com** [https://case.edu/utech/help/knowledge-base/box/box-information](https://case.edu/utech/help/knowledge-base/box/box-information)
- **Qualtrics Information** [https://research.case.edu/corefacilities/CoreFacilityDetail.cfm?n=77](https://research.case.edu/corefacilities/CoreFacilityDetail.cfm?n=77)
- **Cal Frye, Compliance Technologist - cxf244@case.edu**
- **Utech Data Security Information** [https://case.edu/utech/departments/information-security](https://case.edu/utech/departments/information-security)