Research Ethics: A Brief Introduction

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Why Does Research Ethics Matter?

The Immortal Life of Henrietta Lacks

Tearoom Trade: Impersonal Sex in Public Places

Tuskegee syphilis experiment 1932-1972
Objectives

- Apply the three core ethics principles to your own research studies
- Anticipate ethical issues in research
- Identify the risks of participating in research
- Identify the components of informed consent
- Know where to ask for help
Tri Council Policy Statement: *Ethical Conduct for Research Involving Humans* (TCPS 2)

- Overarching Canadian policy framework for research involving human participants
- TCPS published in 1998
- Revised in 2010 and 2014 after extended periods of consultation with relevant stakeholders
- Clear efforts to make TCPS more relevant/applicable to social scientists
TCPS2- Key Principles

Respect for Persons

Welfare

Justice
Centering on the Human Participant

- Who’s Participating?
- Recruitment process
- The consent or assent relationship
- Methods employed
- Risk/benefit assessment
- Confidentiality and privacy protections
TCPS2 - Key Principles

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Principles of Consent

- Free
- Informed
- Ongoing
The Consent Process

- Consent is not a single act of someone agreeing to participate in a study. It is a process that begins with the recruitment and screening of participants and continues throughout their involvement in the research.

- Components of the Consent Process are:
  - Recruitment
  - Information
  - Documenting Consent
  - Debriefing (when required)
Keeping Consent Free

- When someone consents to be a participant in a study it is important that they have done so entirely of their own free will.
- Are there ways in which voluntariness can be compromised?
  - Incentives
  - Authority/Power Relationships
  - Undue Influence
Elements of Informed Consent

Consent form should be written in plain and simple language and provide an accurate description of:

- The research purpose and proposed use of data collected
- Foreseeable risks and potential benefits
- Extent of confidentiality promised
- Requirements of the study (e.g. duration, frequency, nature of tasks and/or measures)
- Withdrawal procedure
- Extent of incentives
- Data management (access and safeguards)
- Possible commercialization
- Disclose any conflicts of interest
- Plans for provision of new information
- Plans for dissemination of results
- Contact information of the researcher in charge of the study
- Contact information of the REB
Conflicts of Interest

Conflict of Interest or even the appearance of a conflict of interest can damage the trust relationships the research community depends on – between REBs, researchers, participants, sponsors, institutions, professional associations, and society.

If you were a participant in a study comparing the user-friendliness of two different but similar computer programs and you heard later that the researcher had developed or invested money in one of them, would you trust the outcome of the study?
Some research requires participants who may lack the necessary cognitive capacity to decide whether or not to participate in research.

In research, capacity refers to the ability of prospective participants to:

- appreciate the possible impact of foreseeable risks and potential benefits on their own well-being
- understand how the conditions of the research may affect them (e.g. time required, difficulty of tasks)
- evaluate whether participation in a particular research project may or may not be in their own best interest
- when research involves those who are not capable of giving consent on their own behalf, researchers must seek the consent of an authorized third party
What scenarios/participant populations can you think of where capacity to consent may be an issue?

- children whose capacity is still developing
- Adults whose capacity is diminishing or fluctuating due to illness or injury
- Those whose capacity remains only partially developed
Assent & Dissent

- Even when an individual’s authorized third party gives consent, it is important to involve the individual to the greatest extent possible.

- As prospective participants they may agree (assent) or not agree (dissent) with their authorized third party’s decision to consent.
Secondary Use of Information

- Information that was collected for one purpose and is now being used to answer the current research question

- Examples of Secondary Use of Information
  - Data from a previous research study
  - Health data
  - Previous student work
  - Other Administrative data

- It matters if data is identifiable or de-identified.
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Types of Risk

- Physical
- Psychological
- Economic
- Social
- Legal
Probability & Magnitude of Risk

- **PROBABILITY**: How likely is it that any participant will suffer any harm as a result of the study?

- **MAGNITUDE**: How severe could the harm be?
Minimal Risk Research

“Research in which the probability and magnitude of possible harms implied by participation in the research is no greater than those encountered by participants in the aspects of their everyday life that relate to the research.”
How To Apply The Principle of Concern for Welfare

- Consider potential impacts on participants’ physical and mental health, on their social or economic circumstances, and on their privacy.

- Consult any groups that may be affected by the research to assess the risk of negative impacts such as stigmatization and discrimination.

- Eliminate and/or minimize risks.

- Maximize benefits.

- Provide accurate and accessible information.
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Understanding Vulnerable Circumstances

- In applying the principle of Justice to research design and review, it is important for researchers to understand vulnerable circumstances from the perspective of the participant.

- No longer categorize participants as “Vulnerable Populations”

- Justification for inclusion and exclusion from participation. These criteria must be relevant to the research question.
How To Apply The Principle of Justice

- In research design it is important to address the following issues:
  - Who are the participants? Why this group and not others?
  - Are any participant groups over- or under-represented because of their vulnerable circumstances?
  - Are there measures in place to treat people in vulnerable circumstances justly in the context of the research?
  - Is there an imbalance of power between participants and researchers?
Centering on the Human Participant

- Who’s Participating?
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- Confidentiality and privacy protections
Confidentiality

The ethical duty of confidentiality refers to the obligation of an individual or organization to safeguard entrusted information. The ethical duty of confidentiality includes obligations to protect information from unauthorized access, use, disclosure, modification, loss or theft. Fulfilling the ethical duty of confidentiality is essential to the trust relationship between researcher and participant, and to the integrity of the research project.
Major Concepts: Confidentiality and Anonymity

- Directly Identifying Information
- Indirectly Identifying Information
- Coded Information
- Anonymized Information
- Anonymous Information
Videos, Photography and Audio Recordings

- Identifiable information

- Recommended to provide participants with a choice to be video recorded, photographed, audio recorded.

- Questions to consider:
  - Are these videos, photographs, audio recordings necessary for the research? Give option to choose.
  - What are the videos, photos, audio recordings going to be used for (analysis only, presentations, publications)?
  - Will images be blurred or voice distorted?
Dissemination

- Anonymize information as much as possible if you have not had participants consent to being directly identified.

- Indirectly identifying individuals should be avoided unless you have obtained prior permission to do so.

- Dissemination can be made in many different formats (blogs, journal articles, presentations etc.) – what matters is if the information could identify your participants and what did they give consent to.
Data Storage/Access

- Data Management Plan

- Data Storage
  - Long-Term – SFU RADAR - [http://researchdata.sfu.ca/node/6](http://researchdata.sfu.ca/node/6)
Review Process

PI submits Application through ORE Application Database

Initial Review for Completeness Conducted by ORE Staff

Faculty Supervisor Reviews and Approves PI’s submission

PI Receives and Responds to Queries

Above Minimal Risk??

ADORE/DORE Review

Minimal Risk

Letter of Approval Issued

Full REB Review

Principal Investigator (PI) / Faculty Supervisor

Office of Research Ethics (ORE)

SFU Research Ethics Board (REB)
Funding Titles and ORE Application Titles

Grant proposals that have been peer-reviewed and funded by granting agencies. Read more

Study Submission Deadlines for Full Board Applications

SFU Chosen as BCEHI Coordinator

The Michael Smith Foundation for Health Research (MSFHR) has awarded SFU the role of project coordination for the BC Ethics Harmonization Initiative for one year. Read more
What to Submit

- A detailed description of the study (Study Details, protocol, proposal, grant application, etc.)
- Consent documents
- Recruitment materials (posters, letters of introduction, email scripts, etc.)
- Any research instruments (questionnaires, interview questions, etc.)
- After you submit, you will not be able to go back to the Database to make changes! Send an email to dore@sfu.ca with the changes (e.g. documents attached) and ORE staff will attach to your application.
What the REB and ORE Look For

Participant Population
- Who? Where from? Vulnerability of participants

Consent process
- Is consent process plan appropriate for the participant population?

Adequacy of consent process
- Study procedures, voluntary participation, right to withdraw, risks, benefits, confidentiality, plan for incidental findings, if applicable

Risks to participants
- What risks does research expose participants to?

Confidentiality
- If/how confidentiality will be maintained?
General Advice

- **Consistency**
  - Make sure details provided are consistent throughout application: recruitment, procedures, consent & assent documents, advertisements, for example.

- **Clarity**
  - Explain your procedures as clearly as possible: what you plan to do, how you plan to do it, to whom & how often.

- **Detail**
  - Complete all sections of the application.
  - Small details are important but balance between providing too much & too little.
Processes for the ethical review of multi-jurisdictional research at SFU are in place:
1. The BC Ethics Harmonization Initiative (BCEHI)
2. Previously reviewed and approved, minimal risk research

These processes take more time so be prepared!

http://www.sfu.ca/ore/bc-ethics-harmonization-initiative.html
Summary

- The 3 **core ethics** principles are the foundation for the ethical reasoning that guides interactions with participants in all research involving human participants.

- Researchers are obligated to provide as much information as individuals need to make a **fully informed choice** to consent or refuse to participate.

- **Context matters** when determining risk to participants and their vulnerability.

- Be **transparent, clear** and **consistent** when describing your research, procedures and risk to the REB and to participants.
Need Help? Let us know!!!

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