



FRIENDS RESEARCH INSTITUTE, INC.
INSTITUTIONAL REVIEW BOARD
1040 Park Avenue, Suite 103
Baltimore, Maryland 21201
410-837-3977 (phone) 410-752-4218 (fax)

Application for Behavioral Study Review

IRB #:

I. PROJECT INFORMATION
1. Project Title:
2. Principal Investigator:
Address:
Telephone Number: Fax Number:
E-Mail Address:
3. Co-Investigator:
Address:
Telephone Number: Fax Number:
E-Mail Address:
4. Primary Contact Person:
Address:
Telephone Number: Fax Number:
E-Mail Address:
5. Any Additional Co-Investigators? [] Yes [] No If yes, please attach additional contact information

II. INVESTIGATOR'S ASSURANCE
The Principal Investigator must assure the Board that all procedures performed under the project will be conducted in strict accordance with all applicable Federal, State and Local regulations and laws regarding the protection of human subjects/participants in research including, but not limited to:
1. Use of qualified personnel to conduct the project according to the protocol approved by the IRB.
2. Ensuring that no changes are made to the approved protocol or consent form without prior IRB approval (except in an emergency to safeguard the well-being of subjects/participants or others).
3. Using the most current, approved, stamped consent form to obtain informed consent from subjects/participants or their legally responsible representative.
4. Prompt reporting of serious adverse events (SAEs) and unanticipated problems to the IRB in writing within 48 hours of discovery, and 24 hours for deaths. For greater than minimal risk studies, all SAEs and unanticipated problems need to be reported. For minimal risk studies, only SAEs and unanticipated problems which are possibly or definitely related to the research study need to be reported.
5. If I will be unavailable to direct this research personally, as when on leave or vacation, I will arrange for a co-investigator to be directly responsible for this study in my absence.
Signature of Principal Investigator Date

III. FUNDING
Funding Source:

Contract or Grant Title:	
Contract or Grant #:	

IV. STUDY SITE

Specify the location where the research will be conducted. *Note: If any facilities are indicated that are not associated with FRI, a letter of support is required, and IRB approval from that institution may be required.*

V. PROTOCOL SUMMARY

Please fill out the information requested in the following categories or cite page numbers if a separate study protocol is attached. If the item does not apply to your research, simply indicate that the question is not applicable.

1. Specific Aims: List the broad, long-term objectives and what the specific research proposed in this application is intended to accomplish.
2. Background and Significance: Provide a brief sketch of the background leading to the present application.
3. Preliminary Studies: Provide an account of the principal investigator's preliminary studies pertinent to the application and a list of all literature cited.
4. Research Design and Methods: Provide a description of the research design and the procedures to be used to accomplish the specific aims of the project.

VI. STUDY POPULATION

1. Describe the characteristics of the subject/participant population such as the anticipated number, age range, gender, ethnic background and health status. Provide a candid discussion of potential problems related to the subject/participant population.
2. Inclusion of Women: Address the inclusion of women unless there is a compelling rationale for proposed exclusion of any sex/gender group.
3. Inclusion of Minorities: Address the inclusion of minorities unless there is a compelling rationale for proposed exclusion of any racial/ethnic group.
4. Inclusion of Children: Address the inclusion of children, if applicable. *{Children are defined in the HHS regulations as "persons who have not attained the legal age for consent to treatments or procedures involved in the research, under the applicable law of the jurisdiction in which the research will be conducted" [45 CFR 46.402(a)].}* (For research involving children, please complete and include the Supplementary Application for Research Involving Children.)
5. Inclusion of Prisoners: Address the inclusion of prisoners, if applicable. *{Prisoner is defined in the HHS regulations as "any individual involuntarily confined or detained in a penal institution. The term is intended to encompass individuals sentenced to such an institution under a criminal or civil statute, individuals detained in other facilities by virtue of statutes or commitment procedures which provide alternatives to criminal prosecution or incarceration in a penal institution, and individuals detained pending arraignment, trial, or sentencing" [45 CFR 46.303(c)].}* (For research involving prisoners, please complete and include the Supplementary Application for Research Involving Prisoners.)
6. Decisionally Impaired: Address the inclusion of this group if there is a clear/compelling reason to include them.

VII. RECRUITMENT

What methods will be used to identify and recruit potential subjects/participants? Attach a copy of all planned

advertisements, flyers, and letters, etc. targeted at or to be sent to potential subjects/participants.

VIII. INFORMED CONSENT

1. Please identify by name, degree, and training the individual(s) who will be authorized to describe the research to subjects/participants, or their representatives, and to administer the consent process. *The personnel administering the consent must have appropriate training and background to insure that subjects/participants give complete and truly informed consent, and to be able to answer any questions regarding the research that subjects/participants may have.*
2. Will all subjects/participants have the capacity to give informed consent? If not, describe the likely range of impairment and explain how, and by whom, their capacity to consent will be determined.
3. Will any information about the research purpose and design be withheld from subjects/participants? If so, please explain the non-disclosure and describe plans for post-study debriefing.

IX. RISK/BENEFIT ASSESSMENT

Note: The potential benefits of the research must justify the risks to human subjects/participants. The risk/benefit ratio of the research must be at least as favorable for the subjects/participants as that presented by standard treatments for their condition. When comparing the risk/benefit ratio of research with that of available alternatives, the alternative of doing nothing should be included in the analysis.

1. Describe the potential benefits, if any, the subjects/participants may receive as a result of their participation in the research and what benefits to society, if any, may be expected.
2. What therapeutic alternative(s) are reasonably available to potential subjects/participants should they choose not to participate in the study? *(These may be research or non-research-based alternatives.)*
3. Describe any potential risks or likely adverse effects subjects/participants may encounter in the study. State the potential risks – physical, psychological, social, legal or other – connected with the proposed procedures and assess their likelihood and seriousness. *(These must be included in the consent form.)*
4. Describe the procedures for minimizing any potential risks.
5. Data and Safety Monitoring Board/Plan: Provide a general description of a monitoring plan that establishes the overall framework for data/adverse event monitoring.
6. Risk Classification: Please check the level* of risk associated with this study.
**According to HHS/FDA regulations "minimal risk" means: "The probability and magnitude of harm or discomfort anticipated in the research are not greater in and of themselves than those ordinarily encountered in daily life or during the performance of routine physical or psychological examinations or tests." When the risks that are associated with a new procedure or product are unknown, they cannot be classified as minimal.*
 minimal greater than minimal unknown

X. DATA COLLECTION & MANAGEMENT

1. How will the data be collected and recorded? If your study uses surveys, questionnaires, or psychological tests, please describe the provisions for administering these measures, the mode of administration, the setting, and if special training or qualifications are necessary.
2. How will the data be coded to protect confidentiality and personal privacy?
3. How will the data be stored during the study?
4. Who will have access to the data and the data codes? If data with subject/participant identifiers will be released, specify

the person(s) and agencies to whom this information will be released.
5. What will happen to the data when the study is completed?

XI. FINANCIAL CONSIDERATIONS
1. Describe all plans to compensate subjects/participants, including provision of services, and other reimbursements, and the conditions that participants must fulfill to receive full or partial payment.
2. Will subjects/participants have to pay for any of the tests or treatments that they receive as part of the research? <i>(Please clarify who will pay for the procedures associated with the study as well as procedures that may be part of standard clinical care. Clarify that insurance and other third party payers may not cover standard procedures if they are associated with a research project.)</i>

XII. REQUIRED DOCUMENTS	
<input type="checkbox"/>	1. Application for Behavioral Study Review
<input type="checkbox"/>	2. Informed consent form(s)
<input type="checkbox"/>	3. Detailed research protocol
<input type="checkbox"/>	4. CV/Biographical sketch
<input type="checkbox"/>	5. Certification of education in research ethics and the protection of human subjects
<input type="checkbox"/>	6. Financial Disclosure form
If Applicable:	
<input type="checkbox"/>	7. If special populations are involved: <ul style="list-style-type: none"> • Supplementary Application for Research Involving Prisoners • Supplementary Application for Research Involving Children • Supplementary Application for Research Involving Pregnant Women, Human Fetuses, and Neonates
<input type="checkbox"/>	8. Assent/parental permission forms
<input type="checkbox"/>	9. If eligible for expedited review, fill out the Application for Expedited Review
<input type="checkbox"/>	10. All surveys, questionnaires, etc. that are indicated in the protocol
<input type="checkbox"/>	11. Subject/Participant recruitment materials, if used (e.g., flyers, advertisements, copy of radio ads)
<input type="checkbox"/>	12. If outside facilities or agencies are used as research sites: letters of agreement (on the facility's letterhead). If these facilities have an IRB, include a copy of the letter of approval for this study.