**ECHN Institutional Review Committee**

**#1 – Protocol Application for the Involvement of Human Participants in Research**

**(Human Subject Intervention)**

Manchester Memorial Hospital, 71 Haynes Street, Manchester, CT 06040 (860) 646-1222, ext. 2084

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**SECTION I: General Information**

**Study Title**: (The title should accurately reflect the purpose of the study. Provide the complete full-length title for the study)

**Study Objective (2-3 sentence summary of study):**

**Principal Investigator Information:**

Name (First, Last, Degree):

Department:

Mailing Address:

Phone #:

E-mail address:

**Co-Investigator(s) Information:** (add additional co--investigators, if necessary)

Name (First, Last, Degree):

Department:

Mailing Address:

Phone #:

E-mail address:

**Student/Resident Researcher(s) Information:** (add additional student researchers, as necessary)

Name (First, Last, Degree):

Department:

Mailing Address:

Phone #:

E-mail address:

**Other study personnel:**

Name (First, Last, Degree):

Department:

Mailing Address:

Phone #:

E-mail address:

**Section II: Collaborating Institutions/Facilities and Other IRB Reviews**

Will the research be conducted only at ECHN? [ ]  **Yes** [ ]  **No (If yes, skip to section III)**

**Other Collaborating Institutions/Facilities**

If you are collaborating with other sites, provide the name of each institution/facility (e.g., other university, nursing home, etc.) and describe the type of involvement of each institution (e.g., recruitment, enrollment/consenting, study procedures, follow-up, data analysis). Indicate if IRC approval/site permission is attached (indicate yes, no or pending). You will need to obtain IRC approval from every collaborating institution that has an IRC before you can initiate research there. Before initiating the research protocol here, the ECHN IRC may request copies of the other site(s) IRC approvals.

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| --- | --- | --- |
| **Name of Institution** | **Describe Involvement** | **IRB Approval/Site Permission Attached?** |
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If the PI has an affiliation/appointment with an Institution listed above, please explain:

Provide additional comments as needed:

**SECTION III: Funding**

It is the responsibility of the Principal Investigator to notify the IRC via an Amendment or at Re-Approval, if the funding source changes in any way.

|  |  |  |
| --- | --- | --- |
| **Funding Source:** |[ ]  Departmental Funds |[ ]  Investigator out-of-pocket |
|  |[ ]  External – Please specify below:  |[ ]  Unfunded |
|  |[ ]  Faculty Grants (Large/Small) |[ ]  Other – Please specify below |
|  |[ ]  Graduate School Award |[ ]   |
|  |[ ]  Undergraduate Research Award |[ ]   |

Provide additional comments as needed:

**For Externally Funded Studies:**

If the research is supported either in whole or in part by external funds (federal, state or private), one COMPLETE copy of each grant application or contract must be included with this application.

**For each funding source, please identify the following:**

**NOTE**: If the PI on the grant/contract is not the PI on this IRC protocol, submit an e-mail with this application in which the PI who is receiving the grant acknowledges use of this protocol under the grant**.**

|  |  |
| --- | --- |
| Name of Funding Source (if ECHN is the recipient of a sub award, list the institution providing the funding then list the primary source of funds): |       |
| Principal Investigator of Contract/Grant**:** |       |
| Contract/Grant Title:(if different from protocol title) |       |
| Grant/Contract Status:(i.e., pending/awarded)(i.e., pending, awarded) |       |

Provide additional comments as needed:

**SECTION IV: Conflict of Interest (IRC will make a determination if this form is not required)**

At the time of protocol submission to the IRC, all investigators are required to review the ECHN Conflict of Interest Policy and complete the attached Questionnaire. Policy and Questionnaire available at <http://www.echn.org/academic-affairs>

For certain studies, this requirement may be waived. For more information, contact pdombek@echn.org

**SECTION V: Human Participants**

**How many participants will be enrolled?**

If you are enrolling more than one population describe the total enrollment for each. Note: Participants are generally considered to be ‘enrolled’ when they sign the consent form or have gone through an oral consent process. Therefore, be sure to account for attrition in your enrollment number.

**If applicable, how many potential participants will be screened?**

When screening procedures are conducted as part of the consent process, participants that fail to screen will be counted as being enrolled in the study.

**Participant Population(s):**

Describe the participant population(s) including gender, ethnicity, and age range.

**Recruitment:**

Describe the recruitment process including *who* will recruit, *when* and *where* recruitment will take place and *how* participants will be identified and recruited (e.g., direct recruitment by study team in person, on the phone, by mail/email/internet, random sampling, referrals from other participants, snowball sampling and/or healthcare providers). Attach copies of all advertisement/recruitment materials for IRB review including phone scripts, web postings, and newspaper advertisements. If recruiting at off-ECHN facility sites, written permission and/or local IRC approval may be required.

**Special Population(s):**

Identify any special participant population(s) that you will be **specifically targeting** for the study.

|  |  |  |
| --- | --- | --- |
| Check **all** that apply  |[ ]  Minors |[ ]  Economically/Educationally Disadvantaged  |
|  |[ ]  Prisoners |[ ]  Members of the Armed Forces |
|  |[ ]  Pregnant Women/Neonates |[ ]  Non-English Speaking |
|  |[ ]  Decisionally Impaired |[ ]  Individuals Living with AIDS/HIV |
|  |[ ]  Employees |[ ]  Other (Please identify): |

**Employees:**

Are you recruiting employees who report to you? [ ]  **Yes** [ ]  **No**

If ‘Yes,” explain why this population is necessary to the study:

**SECTION VI: Drugs/Devices, Genetic Testing, Radiation and Biological Samples**

**Drug/Device Use**

Does the study involve the use of any of the following (check all that apply)?

* An FDA approved drug or medical device [ ]  **Yes** [ ]  **No**
* An investigative/unapproved drug or medical device [ ]  **Yes** [ ]  **No**
* A non-medical device [ ]  **Yes** [ ]  **No**
* A proprietary product [ ]  **Yes** [ ]  **No**
* A biological agent [ ]  **Yes** [ ]  **No**

**SECTION VII: Research Plan**

# Purpose

State the reason for the study, the research hypothesis, and the goals of the proposed study as related to the research question(s).

**Introduction**

State the background information that led to the plan for this project. Please provide references as appropriate and any other related research previously conducted, if applicable.

# Design, Procedures, Materials and Methods

Describe the study design, including the sequence and timing of all study procedures. Indicate expected start and completion dates. Include screening procedures, if any. The IRC strongly suggests that investigators incorporate flexibility into the study design to accommodate anticipated events (i.e. explain how missed study appointments can be made up by participants). **IF A WAIVER OF AUTHORIZATION IS REQUESTED, PLEASE EXPLAIN THE REASONS.** If applicable, describe the use of audiotape and/or videotape and provide justification for use. **If the study includes measures, survey instruments and questionnaires**, identify each and, if available, provide references for the measures. Describe what they intend to measure (relate to purpose/hypothesis) and their psychometric properties (e.g., reliability and validity). Identify any that were specifically created for the study.

### Data Analysis/Justification of Sample Size

For all studies, explain how the data will be analyzed. For qualitative and pilot studies, describe how the proposed sample size is appropriate for achieving the anticipated results. For quantitative studies, provide a power analysis and references for how the sample size was determined. Explain rate of attrition.

### Inclusion/Exclusion Criteria

### List the major inclusion and exclusion criteria. Any proposed exclusion based on gender (women of childbearing potential), age, or race must include justification for the exclusion. Describe the conditions under which participants may be removed from the study, i.e., noncompliance with study rules, determined to be in the best interest of the participant by the P.I., study termination, etc.

# Risks and Inconveniences

Describe the potential risks to participants (and secondary participants, if applicable) and steps taken to minimize risks. Assess the likelihood of the risk occurring and, if it were to occur, the seriousness to the participant. Types of risks to consider include: physical, psychological, social, legal, employment, and financial. Also describe any anticipated inconveniences the participants may experience (time, abstention from food, etc.).

# Benefits

Describe anticipated benefits to the individual participants. If individual participants may not benefit directly, state so here. Describe anticipated benefits to society (i.e., added knowledge to the field of study) or a specific class of individuals. Do not include compensation or earned course credits in this section.

**Risk/Benefit Analysis**

Describe the ratio of risks to benefits. Risks to research participants should be justified by the anticipated benefits to the participants or society. Provide your assessment of anticipated risks to participants and steps taken to minimize these risks, balanced against anticipated benefits to the individual or to society.

# Economic Considerations

Describe any costs to the participants or amount and method of compensation that will be given to them. Describe how you arrived at the amount and the plan for compensation; if it will be prorated, please provide the breakdown. Experimental or extra course credit should be considered an economic consideration and included in this section.

**Data Safety Monitoring**

This is a prospective plan set up by the study investigators to assure that adverse events occurring during studies are identified, evaluated, and communicated to the IRC in a timely manner. Although the investigators initially propose a Data Safety Monitoring **Plan** (DSMP), the IRC must approve the plan and may require revision of the plan. A DSMP is required for all human studies at ECHN except for studies determined to be exempt from continuing IRC review. For studies that present more than minimal risk to participants, the IRC will review and determine on a case-by-case basis whether a data safety monitoring **Board** is most appropriate.

Issues that should be addressed in the DSMP include the following:

1. frequency of the monitoring
2. who will conduct the monitoring (Under ECHN policy a student/resident cannot be the sole person responsible for monitoring the data and safety of the protocol procedures. )
3. what data will be monitored
4. how the data will be evaluated for problems
5. what actions will be taken upon the occurrence of specific events or end points
6. who will communicate to the IRC and how communication will occur

Sample response to issues listed above for minimal risk/slight increase over minimal risk – “Survey results will be monitored by the PI in conjunction with the student investigator once every two weeks (items 1, 2 and 3). Survey responses will be reviewed to monitor for clarity (i.e., the same question is skipped by 5 or more participants). In that case, the question will be revised and an amendment will be submitted to the IRC (items 4, 5 and 6).”

**SECTION VIII: Confidentiality, Data Management & Storage**

If you are going to collect **identified information** please address the following points:

* Describe adequate steps to preserve the confidentiality of the data collected (e.g., hard copy data will be stored in locked file cabinets within a locked office at ECHN.) A master list of codes and identities will be maintained in a secure location apart from research record (coded data), research file and its contents will only be labeled with a code. If linked or coded information is collected, you need to describe the process for coding.

For electronic data specify where data will be kept or stored (e.g., on standalone desktop computer not connected to network/internet), or will be stored on an encrypted mobile computing device, electronic data will only be labeled with a code and/or electronic data will be stored on password protected pc/mc.

If data is stored in a portable device: describe the portable device(s) to be used (e.g., laptop, PDA, portable hard drive including flash drives). Specify whether subject identifiable data will be stored on the device. If so, justify why it is necessary to store subject identifiers on the device.

* Specify who will have access to the data. If there is a code key, specify who on the research team will hold the key, and who will have access to the key.
* Specify if, how and at what point identifiable information will be separated from data.
* Specify how long data will be retained at the conclusion of the study (3 years minimum for majority of research records, 6 year minimum for HIPAA documents).
* If identifiable, sensitive information (illegal drug use, criminal activity, etc.) will be collected, state whether or not a Certificate of Confidentiality was requested and obtained from the National Institute of Health and justify why that decision was made. If a Certificate of Confidentiality was obtained, attach a copy of the Certificate to this application.
* Specify whether any limits to confidentiality exist and identify any external agencies (study sponsor, FDA, IRC, etc.) that will have access to the data for monitoring/auditing purposes.

**Section IX. Timetable**

Provide a detailed timetable scheduling all aspects of the research.

*It is convenient for the reviewer to see the events of the study schedule or duration in the form of a flow chart.* You may reference an attached flow diagram, including expected start and completion dates, and/or describe the time table here: **For example:**

**“Study Title”**

**Major Research Activities**

**Start Date** **End Date**

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|  | ***Jan*** | ***Feb*** | ***Mar*** | ***Apr*** | ***May*** | ***Jun*** | ***Jul*** | ***Aug*** | ***Sept*** | ***Oct***  | ***Nov*** | ***Dec*** |
| ***IRB Approval*** |[ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ]
| ***Screening******Enrollment*** |[ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ]
| ***Intervention*** |[ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ]
| ***Data Analysis***  |[ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ]
| ***Report Preparation*** |[ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ]
| ***Dissemination***  |[ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ] [ ]

**SECTION X: Informed Consent**

**As PI, you are responsible for taking reasonable steps to assure that the participants in this study are fully informed about and understand the study. Even if you are not targeting participants from “Special Populations” as listed on page 3, such populations may be included in recruitment efforts. Please keep this in mind as you design the Consent Process and provide the information requested in this section.**

**Consent Setting**

Describe the consent process including who will obtain consent, where and when will it be obtained, assessment of the patient’s capacity to consent, conditions under which consent will be obtained, any steps to minimize undue influence and any steps to enhance the patient’s independent decision-making, such as a waiting period. Describe how the privacy of the participants will be maintained throughout the consent process. State whether an assessment of consent materials will be conducted to assure that participants understand the information (may be warranted in studies with complicated study procedures, those that require extensive time commitments or those that expose participants to greater than minimal risk).

**Capacity to Consent**

Describe how the capacity to consent will be assessed for participants with limited decision-making capacity, language barriers or hearing difficulty. If a participant is incapable of providing consent, you will need to obtain consent from the participant’s legal guardian.

**Parent/Guardian Permission and Assent**

If enrolling children, state how many parents/guardians will provide permission, whether the child’s assent will be obtained and if assent will be written or oral. Provide a copy of the script to be used if oral assent will be obtained.

**Documentation of Consent**

Specify the forms that will be used for each participant population, i.e., adult consent form, surrogate consent form, child assent form (written form or oral script) or an information sheet. Copies of all forms should be attached to this application in the same format that they will be given to participants.

**Waiver or Alteration of Consent**

The IRB may waive or alter the elements of consent in some minimal risks studies. If you plan to request either 1) a **waiver of consent** (i.e., participants are not aware of their participation in the study and will not be asked to give consent), or 2) a **waiver of signed consent** (i.e., participants participation in the study after reading a study information sheet implies consent; however, no signed consent is obtained), please answer the following questions using specific information from the study:

Please complete all questions in either #1 or # 2 (as applicable):

**1) Waiver of consent:**

* Why is the study considered to be minimal risk?

* How will the waiver affect the participants’ rights and welfare? The IRC must find that participants’ rights are not adversely affected.
* Why would the research be impracticable without the waiver?

* How will important information be returned to the participants, if appropriate?

**2) Waiver of signed consent:**

* Why is the study considered to be minimal risk?

* Does a breach of confidentiality constitute the principal risk to participants?

* Would the signed consent form be the only record linking the participant to the research?

* Does the research include any activities that would require signed consent in a non-research setting? For example, in non-research settings, normally there is no requirement for written consent for completion of questionnaires.

**HIPAA Authorization**

At ECHN, the following sites are covered entities under the Health Insurance Portability and Accountability Act:

1. Manchester Memorial Hospital

2. Rockville General Hospital

3. Woodlake at Tolland

4. ECHN Family Health Care Center

If research participants are recruited through these entities:

* Obtain an Authorization to use and disclose Protected Health Information (PHI), or
* Request Waiver of Authorization. To request a waiver, the following elements must be satisfied:

|  |
| --- |
| 1. Use or disclosure involves no more than minimal risk to the privacy of the individuals, based on the presence  of at least the following elements:a. an adequate plan to protect health information identifiers from improper use or disclosure,b. an adequate plan to destroy identifiers at the earliest opportunity absent a health or research justification or legal requirement to retain them, andc. adequate written assurances that the PHI will not be used or disclosed to a third party except as required by law, for authorized oversight of the research study, or for other research uses and disclosures permitted by the private rule. |
| 2. Research could not practicably be conducted without the waiver or alteration; and |
| 3. Research could not practicably be conducted without access to and use of PHI. |

A "Partial Waiver" may be sought and granted during the recruitment/screening phase of a study. If it is then determined that the participant is eligible for the study, an authorization would have to be obtained prior to or at the first visit.

**Section XI: References and Literature:**

**Required Attachments:**

1. The original study protocol (if applicable) with this completed ECHN Protocol Submission form.
2. Copy of the approval letter from any external IRB (if applicable, e.g. UNECOM, CCMC, UCONN)
3. Informed Consent form
4. Any Recruitment Materials
5. Letter of support from Department Chair/Director
6. Principal Investigator’s CV
7. CITI training or NIH training certificate
8. Authorization to Use or Disclose PHI OR Waiver of Authorization (if applicable)

**Principal Investigator Certification**

I understand Eastern Connecticut Health Network (ECHN) policies concerning research involving human participants and I agree:

1. To comply with all IRC policies, decisions, conditions, and requirements;
2. That this study has been designed, to the best of my knowledge, to protect human participants engaged in research in accordance with the standards set by Eastern Connecticut Health Network, the United States Department of Health and Human Services, the Food and Drug Administration, and any other sponsoring agency;
3. To obtain prior approval from the IRC before amending the research protocol or the approved consent/assent form;
4. To report to the IRC in accordance with IRC policy, any serious adverse event(s) and/or unanticipated problem(s) involving risks to participants;
5. To submit the Re-Approval letter at least yearly, or as needed;
6. That my participation and the participation of any co-investigators does/do not violate Eastern Connecticut Health Network's policy on Individual Conflicts of Interest in Research;
7. That each individual listed as study personnel in this application has a) completed the required human subjects training, and b) are knowledgeable of the study procedures described in the protocol;
8. That each individual listed as study personnel in this application possesses the necessary training and experience for conducting research activities in the role described for them in this research study.

 9. All external researchers are required to sign the ECHN Confidentiality Statement for Researchers ` (available on the ECHN Portal).

Furthermore, by signing below, I also attest that I have appropriate facilities and resources for conducting the study.

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| **Original Signature of Principal Investigator** | **Date** |

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| **Original Signature of Student/Resident Investigator****(Only for Student/Resident-Initiated Research)** | **Date** |

Rev. 6/2012, 11/2012, 12/2012, 5/2013, 11/2014, 12/2014, 1/2016, 6/2016, 8/2017