



FOUNDED BY BRIGHAM AND WOMEN'S HOSPITAL
AND MASSACHUSETTS GENERAL HOSPITAL

| | |
|--------------------------|--|
| Title: | Human-Subject Protection Education and Training Requirements for Investigators and Study Staff |
| Department: | Human Research Affairs/Research Management |
| Policy Type: | <input checked="" type="checkbox"/> Partners System-wide <input type="checkbox"/> Partners System-wide Template <input type="checkbox"/> Partners Corporate <input type="checkbox"/> Partners Corporate Departmental <input type="checkbox"/> Entity |
| Applies to: | Employees, Professional Staff or Other Agents of Brigham and Women's Hospital (BWH), Faulkner Hospital (FH) and Massachusetts General Hospital (MGH) |
| Approved by: | Chief Scientific Officer |
| Approval Date: | September 9, 2010 |
| Effective Date: | September 9, 2010 |
| Revision Date(s): | |
| Next Review Date: | September 9, 2013 |
| Contact Person: | Director, Human Research Review and Compliance |

KEYWORDS:

IRB, Institutional Review Board

PURPOSE:

The purpose of this policy is to ensure that individuals conducting non-exempt human-subjects research overseen by the Partners Human Research Committee (PHRC) understand the ethical principles and regulations related to the protection of human subjects of research.

DEFINITIONS:

See Definition of Human-Subjects Research

POLICY STATEMENT:

The Brigham and Women's Hospital (BWH), Faulkner Hospital (FH) and the Massachusetts General Hospital (MGH) have a legal and ethical responsibility to protect the rights and welfare of human subjects participating in research conducted or sponsored by BWH, FH or MGH or under the auspices of the hospitals, or in which the hospitals are otherwise engaged regardless of the location of the research or source of funding. Consistent with these responsibilities, the BWH, FH and MGH require every individual engaged in non-exempt human-subjects research overseen by the Partners Human Research Committee to complete the web-based

Collaborative IRB Training Initiative (CITI) basic biomedical research education and training module prior to their involvement in the research and the biomedical continuing education program every three years. The PHRC may accept an equivalent human-subject protection education program on a case-by-case basis. Notwithstanding this policy, the PHRC may require an investigator to fulfill additional education and training requirements based on the type of research (e.g., IND/IDE sponsor-investigator research) or as part of remedial education.

PROCEDURES:

1. New non-exempt research involving human subjects will not be approved by the PHRC until all of the study staff listed on the protocol have completed the human-subject protection education requirements (the CITI program) including, when applicable, continuing education requirements. Completion of the CITI education programs will be recorded in the PHRC research community database.
2. The addition of new study staff will not be approved by the PHRC unless the individual(s) being added via amendment has completed the human-subject protection education requirements (the CITI program) including, when applicable, continuing education requirements.
3. At continuing review, the research will not be re-approved by the PHRC unless all of the study staff listed on the protocol have completed the human-subject protection education requirements (the CITI program) including, when applicable, continuing education requirements.
4. The Principal Investigator may elect to remove individuals from the study staff who have not completed the education requirements so that the study may be re-approved; however these individuals may not continue to function as part of the study staff unless and until they have completed the education requirements and an amendment to add them to the study staff has been submitted and approved by the PHRC.
5. Principal Investigators are responsible for ensuring that the study staff listed on their protocols complete their continuing education requirements every three years. Completion of the CITI education requirements can be verified in the protocol record in Insight/eIRB by opening the Staff & Access tab or by searching the BWH/FH/MGH CITI education/training list on the PHRC website. Failure on the part of the study staff to comply with the human-subject protection continuing education requirements will be considered noncompliance with PHRC policies and procedures and should be reported at continuing review as a minor protocol deviation/violation.
6. In addition to the mandatory education and training, investigators and study staff are strongly encouraged to take advantage of the many education and training opportunities offered through the BWH Center for Clinical Investigation (CCI) and MGH Clinical Research Program (CRP).

DEVELOPMENT AND CONSULTATION

Human Research Office

| | | |
|--------------|-----------------------|--------------------------|
| Reviewed by: | Original Review Date: | Revision Approval Dates: |
| | | |
| | | |

| | | |
|--|--|--|
| | | |
| | | |
| | | |
| | | |
| | | |