POL 06002  FHI 360 Standards for Research Involving Human Subjects

PURPOSE:

To define the ethical, regulatory, and quality standards for the conduct of research involving human subjects at FHI 360.

SCOPE:

This policy applies to FHI 360 staff involved in the conduct of research involving human subjects. Additional requirements may be applicable depending on Sponsor agreements/contracts.

DEFINITIONS:

1. **Clinical research** A type of human subject research for which an investigator directly interacts with human subjects or on material of human origin with the goal of understanding human disease and improving human health. It includes the study of disease mechanisms, therapeutic interventions, epidemiology, and clinical trials (US Department of Health and Human Services Public Health Service: Grant Application PHS 398).

2. **Clinical trial** A research study in which one or more human subjects are prospectively assigned to one or more interventions (which may include placebo or other control) to evaluate the effects of the interventions on biomedical or behavioral health-related outcomes. (Source: 45 CFR 46.102(b))

3. **Human subject** A living individual about whom an investigator (whether professional or student) conducting research: (i) Obtains information or biospecimens through intervention or interaction with the individual, and uses, studies, or analyzes the information or biospecimen; or (ii) Obtains, uses, studies, analyzes or generates identifiable private information or identifiable biospecimens (Source: 45 CFR 46.102(e)(1)).

4. **Investigational product (IP)** A pharmaceutical form of an active ingredient or placebo being tested or used as a reference in a clinical trial, including a product with a marketing authorization when used or assembled (formulated or packaged) in a way different from the approved form, or when used for an unapproved indication or when used to gain further information about an approved use. (Source: ICH E6(R2)).

5. **Intervention** Both physical procedures by which information or biospecimens are gathered (e.g., venipuncture) and manipulations of the subject or the subject's environment that are performed for research purposes.

6. **Research** A systematic investigation, including research development, testing and evaluation, designed to develop or contribute to generalizable knowledge. Activities that meet this definition constitute research for purposes of this
policy, whether or not they are conducted or supported under a program that is considered research for other purposes. For example, some demonstration and service programs may include research activities. (Source: 45 CFR 46.102(l)).

7. Sponsor

An individual, company, institution, or organization which takes responsibility for the initiation, management, and/or financing of clinical and non-clinical research involving human subjects.

POLICY:

1. FHI 360 staff shall apply the following ethical, scientific, and regulatory requirements in the design, conduct, recording, and reporting of research involving human subjects. Compliance with these requirements provides assurance that the rights, safety, and well-being of human subjects are protected, and that the data generated are credible.

   1.1. FHI 360 human subject research shall be guided by the ethical principles of the Belmont Report (1979) and the principles that originate from the Declaration of Helsinki.

   1.2. FHI 360 staff shall apply Title 45 Code of Federal Regulations (CFR), Part 46, including subparts A, B, C and D (as applicable) to all federally funded research. The fundamental commitment to the protection of human subjects shall be applied to all human subject research conducted by FHI 360, regardless of funding source or site of the research.

   • Studies conducted under a US Food and Drug Administration (US FDA) Investigational New Drug Application (IND) or Investigational Device Exemption (IDE) shall comply with applicable 21 CFR regulations, including 21 CFR 50 - the US FDA regulations for human subject protections.

   1.3. FHI 360 human subject research activities shall be implemented in compliance with applicable laws and regulations. Where there are discrepancies between local ethical standards, local laws, and regulations, and/or international standards, these will be resolved in consultation with qualified staff in the Office of International Research Ethics (OIRE) and/or Clinical Regulatory and Compliance (CRC) department and/or sponsor (as applicable).

   1.4. Clinical trials shall be conducted according to the standards of the International Council for Harmonization Guideline for Good Clinical Practice (ICH E6(R2).

   1.5. FHI 360 staff are responsible for implementing and maintaining quality assurance and quality control systems with written procedures that define the research activities to ensure that studies are conducted, and data generated, documented (recorded), and reported in compliance with the protocol, and applicable regulatory requirement(s).

   1.6. Human subject research shall be monitored to assess the protection of subjects’ rights and welfare, and to ensure that the study is being conducted in accordance with the protocol and applicable regulatory requirement(s).

   • Deviations from the Institutional Review Board (IRB)/independent ethics committee (IEC) approved protocol must be documented and reported in accordance with FHI 360 policies and standard operating procedures as well as any local IRB/IEC policies.
1.7. FHI 360 staff are responsible for securing agreement from all involved parties to ensure direct access to all study related sites, source data/documents, and reports to allow for monitoring and auditing by FHI 360 qualified staff, the sponsor (when applicable), and inspection by health regulatory authorities.

1.8. Agreements/contracts made with the investigator/institution and any other parties involved with the research study (including sponsors) shall be in writing, as part of the protocol or in a separate agreement.

1.9. Exceptions to this policy may be allowed with written approval from the Business Unit Director, the appropriate research director (or equivalent), Clinical Regulatory and Compliance Director, and sponsor representative (when applicable).
   - Exceptions shall be documented (1) in the study protocol, if appropriate and feasible; and (2) by a memorandum to the study files with attached written approvals.

2. RESPONSIBILITIES:
   2.1. Study team leaders, applicable department research directors and principal investigators conducting human subject research are responsible for designing and implementing research activities in accordance with this policy.
   2.2. FHI 360 staff is responsible for ensuring the implementation of study procedures according to these standards and documenting any deviations according to established procedures.

For policy interpretation or questions please contact: Director, Office of International Research Ethics or Director, Clinical Regulatory and Compliance.

RELATED DOCUMENTS:

1. Policies
   - POL 06003: Communication with Regulatory Authorities Governing Clinical Research
   - POL 06004: Training Requirements for Staff Engaged in Research Involving Human Subjects
   - POL 06005: Protection of Human Subject Committee (PHSC)
   - POL 06006: Criteria for Protection of Human Subjects Committee (PHSC) Approval of Research
   - POL 06007: Institutional Review Board Review for FHI 360 Studies
   - POL 06011: Children as Research Participants
   - POL 06013: Identification, Assessment and Reporting of Unanticipated Problems and other Adverse Events Involving Risks to Human Subjects or Others
   - POL 06014: Research Integrity and Misconduct
   - POL 06015: Written Translation Requirements for FHI 360 Human Subject Research
   - POL 06016: Handling of Investigational and Non-Investigational Products in Human Subject Research Activities
   - POL 06017: Financial Conflict of Interest in Research Activities Funded through US Public Health Service
2. **Standard Operating Procedures**
   - SOP 00001: Management of FHI 360 Policies, Standard Operating Procedures and Associated Documents

3. **Appendices**
   - NA

**REFERENCES:**

**POL 06002**  FHI 360 Standards for Research Involving Human Subjects

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**POLICY REVISION HISTORY:**

<table>
<thead>
<tr>
<th>POL#</th>
<th>Date Reviewed (DD MMM YYYY)</th>
<th>Summary of Changes</th>
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<tbody>
<tr>
<td>118</td>
<td>May 2000</td>
<td>New</td>
</tr>
<tr>
<td>118</td>
<td>November 2002</td>
<td>Changes reflected in 118A</td>
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<td>October 2004</td>
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<tr>
<td>118C</td>
<td>December 2004</td>
<td>Number change to 118.03</td>
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<td>January 2007</td>
<td>Changes reflected in 118.04</td>
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<td>118.04</td>
<td>October 2009</td>
<td>Changes reflected in 118.05</td>
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<tr>
<td>POL 06002</td>
<td>December 2012</td>
<td>New POL# for migration to EDMS</td>
</tr>
<tr>
<td>POL 06002</td>
<td>April 2014</td>
<td>Editorial changes and update to requirements</td>
</tr>
<tr>
<td>POL 06002</td>
<td>August 2015</td>
<td>Minor change to replace references to RAQA with CRC. Corrected related documents with updated titles. Removed a related document since it's obsoleted. Minor template formats.</td>
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<tr>
<td>POL 06002</td>
<td>January 2017</td>
<td>Updated scope, deleted clinical research definition and updated the definitions for clinical trial/study and investigational product. Added definition for sponsor. Updated policy names and added ICH as a reference. Deleted Appendix reference since the Belmont Report is listed as a reference.</td>
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<tr>
<td>POL 06002</td>
<td>21 MAY 2019</td>
<td>Updated definitions to comply with ICH GCP E6(R2) and revised common rule 45 CFR 46.</td>
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<tr>
<td>POL 06002</td>
<td>07 JUL 2021</td>
<td>Updates to definitions and minor edits for clarity.</td>
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REVISION HISTORY

Version 04 Effective on 12-Jul-2019
TI

Version 05 Effective on 27-Aug-2021
Updates to definitions and minor edits for clarity. Collaboration with Sara Tenorio, Julia and Emily Namey
12JUL2021

DOCUMENT ELECTRONIC SIGNATURES

DOCUMENT APPROVAL WORKFLOW

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I am the author of this document.
Signed 1:08:20 PM UTC 16-Jul-2021

Required Workflow Steps for this Category

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I have reviewed and approve this document.
Signed 2:06:59 PM UTC 26-Jul-2021

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I have reviewed and approve this document.
Signed 3:39:02 PM UTC 27-Jul-2021

Policy Approval 1
Patrick Fine
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I have reviewed and approve this document.
Signed 7:54:04 PM UTC 28-Jul-2021

Policy Approval 2