Drexel University Human Research Protection

The Office for Human Research Protections (OHRP) has made an allowance for the Institutional Review Board (IRB) of one institution to act on behalf of the relying institution’s IRB via an IRB Authorization Agreement. The DU IRB refers to this IRB Authorization Agreement as a “Letter of Reliance”. The intent of the agreement is to help minimize or reduce the burdens of review and redundancy in work load when two or more institutions will act together on the same protocol.

When a DU investigator seeks to work collaboratively with another institution, we ask that the following procedures be taken into consideration. Please note that the Letter of Reliance can only be executed with an institution:
- which holds a valid Federal Wide Assurance number (FWA) that is not expired
- whose review was approved at an Expedited or Full Review Level.

If the research project was Exempt from the requirement of an IRB review and approval, the authorization agreement is not required. However, application documents must be submitted to the Drexel University IRB for review.

The process of finalizing the DU IRB Letter of Reliance is executed by a representative from DU Human Research Protection (HRP) with the other institution’s IRB or their Office of Research Compliance.

There are two primary examples of how the process of securing a Letter of Reliance may apply to a DU investigator.

- **Example 1**: DU IRB is the IRB of record if the research is conducted at DU facilities which include DU patients or patient records, students or employees. In this example the other institution will rely on the DU IRB when collaborating with DU faculty.
- **Example 2**: the other institution’s IRB will be the IRB of record if the research will be conducted on their property, with their patients or patient records, or with their students or employees.

Please note that the DU IRB will always reserve the right of determining whether or not an agreement will be executed, as well as the right to review a proposal in addition to, and independent of any other institution’s IRB.

1. Selecting the IRB of record.
   a. **Research conducted at DU facilities**, with its faculty, staff, students, patients or their patient’s records, requires that the DU IRB will be the IRB of record, and will review the proposal independently of another institution’s IRB. The DU IRB will be responsible for oversight of the related compliance activities and reporting requirements in accordance with both OHRP and FDA guidelines. (Should this be the case follow directions in #3 below).
b. Research conducted at another institution, (regarding the examples cited above in 1a), will require the other institution to be responsible for the IRB review, related compliance activities and reporting requirements in accordance with both OHRP and FDA guidelines (Should this be the case follow directions in #4 below).

2. Once the Principal Investigator (PI)* has determined where the research will primarily take place, and who will be involved, the PI must contact their IRB and/or compliance office, as well as the IRB and/or compliance office of the other institution to confirm that such an agreement is acceptable to both IRBs/institutions.

3. If the DU IRB will be the IRB of record, the DU PI must provide the following documentation to the IRB:
   a. A cover memo requesting the reliance agreement including a detailed description of the role(s) the non-DU PI or student/ key personnel will play in the research study at the other institution. This letter should be on official letterhead and signed by the PI and include the name(s) of the student/key personnel.
   b. A complete standard application in accordance with the appropriate directions and IRB policies
   c. If research activities will be conducted at DU and will involve non-DU research personnel, the following documentations must be submitted: proof of DU HIPAA I and II trainings, proof of CITI Human Subjects Research Protection Training(s): Basic and HIPS; HRP 201-Contact Information Form, Financial Interest Disclosure Form(s) for each personnel;
      i. Investigative team members without a DU affiliation, that are affiliated with an institution that has not filed for a Federal Wide Assurance (FWA) with OHRP will be required to complete and sign an “Unaffiliated Investigator Agreement”;
      ii. If the proposal involves use or access to Protected Health Information (PHI), the relying institution needs to be documented in the consent form(s), or any unaffiliated personnel may need to complete the Business Associate Agreement. (Both agreements are available from the DU IRB coordinator)
   d. Relying Institution: The name, title and address of the Authorizing Official that will sign the Authorization Agreement. The contact name and information for any inquiries during the authorization agreement process. The contact information for the IRB/Office of Research Compliance, inclusive of their FWA number and a copy of the institution’s FWA.
e. During the DU IRB review process, the IRB coordinator will contact the other institution directly to confirm their agreement with the terms of an official reliance between the institutions;

f. Once the DU PI receives a letter of approval from the DU IRB it is expected that the PI will provide a copy of all official IRB communications and approval documents to the relying institution’s IRB.

g. Please note: PI’s requesting a “Letter of Reliance” on behalf of a protocol that already has DU IRB approval, will follow the directions above and also submit a modification request to the IRB by use of form HRP-213 “Modification of Approved Human Research.”

4. If the other institution will be the IRB of record, the DU PI needs to provide the following documentation to the DU IRB:
   a. A cover letter from the DU PI stating the request for a reliance agreement including a detailed description of the role(s) the DU PI or student/ key personnel will play in the research study at the other institution. This letter should be on official letterhead and signed by the PI and include the name(s) of the student/key personnel.
   b. Required Trainings: DU HIPAA I and II trainings, CITI Human Subjects Research Protection Training(s): Basic and HIPS; Required Forms: HRP 201-Contact Information Form, Financial Interest Disclosure Form(s) for each personnel;
   c. IRB of Record: The name, title and address of the Authorizing Official that will sign the Authorization Agreement. The contact name and information for any inquiries during the authorization agreement process. The contact information for the IRB/Office of Research Compliance, inclusive of their FWA number and IRB number (IRB that reviewed and approved the study and a copy of the institution’s FWA.
   d. A copy of the initial approval letter and current continuing review approval letter, the currently approved consent form(s), application, advertisements, and any other documents approved by the other institution’s IRB that are in support of the research proposal;
   e. A copy of the modification approval letter indicating the names of the DU personnel added to participate in the research; A copy of the revised consent form adding DU to the participation of the research project, and adding DU to the HIPAA Section.
   f. The DU PI will remain in contact with the DU IRB by providing a copy of all official communications and documents submitted to the other institution’s IRB as long as the protocol remains active. Examples of communications are adverse event reports, continuing reviews/renewals, final reports, amendments to protocols, protocol violations and deviations.
g. It is the PI of record responsibility to remain in contact with the reviewing IRB and to follow any and all directions and procedures they require; this may include additional institution specific training or security clearances.

h. It is strongly recommended that the PI of record maintains open lines of communication with the IRB of record to help assure compliant conduct of research.

5. Once the Letter of Reliance has been finalized the DU PI will receive official notice from the DU IRB that they may initiate their protocol. The DU IRB coordinator will email to the PI's attention the DU IRB approval letter and a copy of the final executed Letter of Reliance.

*Please note DU IRB guidelines will only permit faculty and staff members to serve as PI. Students, medical research residents, fellows, and post-doctoral fellows are always co-investigators.

When submitting your proposal or letter of request for a “Letter of Reliance” to the DU IRB on behalf of a student project, the student will be noted as co-investigator or as key personnel and all documents regarding the study will be addressed from and in the name of the PI. The student will also be listed as noted above.

Any additional questions may be addressed by calling a representative from Human Research Protection at 215-762-3944.

The LOR Submission is to be emailed to HRPP@drexel.edu.