1. **PURPOSE**

1.1. All Institutional Review Board (IRB) communications regarding its review of research activities are in the form of written correspondence. The IRB communicates concerns and suggestions regarding human subject protection issues to Investigators following each step of its review. This policy and procedure describes the process for communicating results to study team and other institutional officials.

2. **POLICY**

2.1. The IRB reports its findings and actions to the Principal Investigator (PI) and study team members via written letter (correspondence) that is part of the electronic project record.

2.2. The IRB reports its findings and actions to the Institutional Official or his/her Designee.

2.3. The IRB reports or makes available its findings and actions to the regulatory agency(ies) when the research is overseen or regulated by those agencies, and they require separate reporting (e.g., to OHRP when the research is covered by DHHS regulations and/or to the FDA when the research is FDA-regulated).

2.4. The study PI and designated contacts will be notified of a review outcome (e.g., approve, approve with modifications required to secure final approval, defer, table, or disapprove for protocol reviews; the resulting determinations of an investigation of non-compliance, suspension or termination of IRB Approval, and unanticipated problem involving risks to subjects or others) via portal-generated e-mail when the correspondence letter has been added to the project record.

2.4.1. These reporting procedures are to be completed within 10 business days of the IRB meeting or receipt of the completed Non-Committee Review materials.

2.4.2. A summary report of the review outcomes will be made via e-mail, phone, or in-person meeting with the PI within 10 business days of the IRB meeting and serves as notification if final correspondence is delayed due to technical issues or other unforeseen circumstances.

2.4.3. When the IRB disapproves research, the IRB will provide the PI with a statement of the reasons for the decision and gives the PI an opportunity to respond in person or in writing.

2.5. Reporting to external or regulatory agencies overseeing the research (e.g., OHRP, FDA, DOD, etc.) of Serious Non-Compliance; Continuing Non-Compliance; Suspension of IRB Approval; Termination of IRB Approval; and Unanticipated Problem Involving Risks to Subjects or Others is to take place within 30 business days from the recognition of a reportable event.

3. **PROCEDURES: IRB Staff**

3.1. After the review has been submitted, the electronic system will provide a letter template that corresponds to the review type and recorded action or determination.
3.2. The IRB Staff will refer to Worksheet: Communication of Review Results and prepare the letter by confirming the appropriate template and generating the letter. The action of generating the letter will integrate the study-specific information with the letter template.

3.3. The IRB Staff will review the draft correspondence for accuracy and correct as needed.

3.4. The IRB Staff will send the letter. This action generates a PDF copy of the correspondence and attaches this to the permanent submission record while sending an e-mail notification to the PI and designated study contacts. This also transitions the electronic record to the final state as determined on the submitted review.

3.5. The IRB Staff makes relevant IRB findings and actions available to offices and committees such as the Sponsored Projects Administration (SPA), the Office of Research Integrity and Safety (ORIS), or the Office of Biosafety.

3.5.1. This information is communicated by allowing guest access to the electronic submission record or by providing copies of materials to relevant people.

3.6. The IRB Staff will report the results of Administrative Review for Determination of Human Subject Research, Developmental Approval, Exempt Determination, and reliance upon an External IRB to the study team via correspondence letter that is part of the electronic project record. The study PI and designated contacts will be notified of the review outcome via system-generated e-mail when the correspondence letter has been added to the project record.

3.7. The IRB will acknowledge a request for closure via correspondence letter that is part of the electronic project records. The study PI and designated contacts will be notified of acknowledgment of protocol closure via system-generated e-mail when the correspondence letter has been added to the project record.

3.8. The electronic letter will have the typed name of the IRB Chair for full board reviews or other IRB staff member for other reviews. The auto-generated letter is considered the official approval from the IRB.

4. MATERIALS

4.1. WORKSHEET: Communication of Review Results
4.2. WORKSHEET: Calculation of Approval Intervals

5. REFERENCES

5.1. 45 CFR 46.103(b)(4)(i)