# UNIVERSITY OF TENNESSEE GRADUATE SCHOOL OF MEDICINE INSTITUTIONAL REVIEW BOARD COMMUNICATION FROM THE IRB

# I. **PURPOSE**

To document the policy and procedures used by University of Tennessee Graduate School of Medicine Institutional Review Board (UTGSM IRB) regarding written communication from the IRB.

### II. SCOPE

This SOP applies to the IRB administrative staff and IRB members.

### **Personnel Responsible:**

UTGSM IRB administrative staff and members

### III. BACKGROUND

The IRB is required to prepare and maintain adequate documentation of its IRB activities. This includes copies of all correspondence between the IRB and investigators.

#### In Accordance With:

45 CFR 46.115 and 21 CFR 50, 56

FDA Guidance for Institutional Review Boards and Clinical Investigators 1998 Update located at <u>http://www.fda.gov/RegulatoryInformation/Guidances/ucm126425.htm</u>

Compliance with this policy also requires compliance with state or local laws or regulations which provide additional protections for human subjects.

# IV. **PROCEDURES**

- 1. Following full board review, expedited review, or determination of exempt status of applications to conduct research, the administrative staff will collect any forms documenting primary and / or secondary reviews, subcommittee or consultant reports, articles or informational documents used in the review.
- 2. The IRB administrative staff will prepare, review and send correspondence concerning IRB review and actions. The IRB will notify investigators of its decision to approve, disapprove, defer, and/or seek modifications of research activity, submissions or informed consent documents within 10 days of the meeting.

- 3. If the IRB defers or disapproves a research activity, written correspondence to the PI will include a statement of the decision and give the investigator an opportunity to respond in writing to these concerns.
- 4. Letters to investigators will include:
  - a. The identification number given to the study by the IRB;
  - b. The protocol title and number (if available), version dates or designations;
  - c. Date and version number of the consent document;
  - d. Date of IRB review and determination;
  - e. Duration of approval and date of continuing review by the IRB;
  - f. If conditionally approved or deferred, a list of provisos that must be met and a statement that the research cannot begin until the investigator receives formal written notification of IRB approval following the response(s) to the provisos.
  - g. If deferred, the basis for the IRB's decision and a statement that the investigator may resubmit the protocol.
  - h. All paper correspondence will be on UTGSM IRB letterhead. An Email from the Chair, Director or designee may satisfy any requirement for written notice.
  - i. A copy of all correspondence will be kept in iMedRIS for the study.