PURPOSE
To describe the policies and procedures for researcher reporting of minor protocol deviations.

REVISIONS FROM PREVIOUS VERSION
None

POLICY
Federal regulations require the IRB to review and approve proposed changes to research studies before initiation of these changes, except when changes are “necessary to eliminate apparent immediate hazards to the subject” [45 CFR 46.103(b)(4)(iii)]. Most proposed changes are reviewed through submission of amendments. Any changes that are made to eliminate apparent immediate hazards to a participant should be reported as soon as possible after they occur as a protocol deviation. Deviations range in seriousness according to how the changes may impact subject safety, the degree of noncompliance with federal and state regulations, and the degree of foreknowledge of the event. Deviations must be reported to the IRB with a description of the deviation, its impact on participant safety (if any) and a description of how similar events will be avoided in the future. Once reported, the IRB can make a decision regarding an appropriate response or remedial action. Remedial actions may involve excluding data that was obtained inappropriately or a recommendation for additional monitoring of study procedures. Note that repeated deviations of the same type may be an indication that an amendment is needed to permanently change study criteria.

Major protocol violations (involve risks)

Accidental or unintentional protocol violations that involve risks to participants or others must be reported to the IRB according to the IRB SOP 401 – Reporting and Review of Events Involving Risks to Participants or Others. A major violation is one that may impact participant safety or alter the risk/benefit ratio, compromise the integrity of the study data, and/or affect participants’ willingness to participate in the study. Major violations require prompt reporting and should be reported to the IRB within 7 calendar days of the researcher’s knowledge of the deviation. Reports should be made using the Event Form.
Minor protocol deviations (involve no risks)

Minor or administrative protocol deviations are defined as a one-time deviation from the IRB-approved protocol that involve no risks to subjects or others. A minor deviation is one that does not impact participant safety, compromise the integrity of the study data, or affect participants’ willingness to participate in the study.

Examples of minor or administrative deviations could include:

- Missing pages of signed consent form;
- Inappropriate documentation of informed consent, including:
  - missing researcher signature and/or date;
  - copy not given to the person signing the form;
  - someone other than the participant dated the consent form;
  - individual obtaining informed consent not listed on IRB approved project team list.
- Use of invalid consent form, i.e., consent form without IRB approval stamp or outdated/expired consent form;
- Failure to follow the approved study procedure that, in the opinion of the PI, does not affect subject safety or data integrity;
  - Study procedure conducted out of sequence;
  - Omitting an approved portion of the protocol;
  - Failure to perform a required lab test;
  - Enrollment of ineligible subject (e.g. participant’s age was 6 months above age limit);
  - Study procedure conducted outside of required timeframe;
- Over-enrollment.

RESPONSIBILITIES
Execution of SOP: Researchers.

PROCEDURE
Researcher Reporting Requirements for Minor Protocol Deviations

1. Members of the research team must immediately report the occurrence of a protocol deviation to the Principal Investigator (PI).
2. All minor deviations should be recorded by the research team and summarized for the IRB at the time of continuing review using the Protocol Deviations Report (available on the OIRB website and IRBNet).
3. Additional reporting requirements to federal agencies, study sponsors, or other entities also may be required as determined by the PI.
4. The IRB may request more stringent requirements for reporting protocol deviations for individual research studies if it is deemed necessary.

REFERENCES

45 CFR 46.103(b)(5)