Title: Review of Protocol Deviations / Violations

SOP Code: SOP 11/V1

Effective Date: aa/bb/cccc

[The IEC members (author/s, reviewer/s) and Chairperson will sign and date the SOP on this first page]

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Table of Contents:

<table>
<thead>
<tr>
<th>No.</th>
<th>Contents</th>
<th>Page No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Purpose</td>
<td>2</td>
</tr>
<tr>
<td>2</td>
<td>Scope</td>
<td>2</td>
</tr>
<tr>
<td>3</td>
<td>Responsibility</td>
<td>2</td>
</tr>
<tr>
<td>4</td>
<td>Detailed Instructions</td>
<td>2</td>
</tr>
<tr>
<td>5</td>
<td>Annexures</td>
<td>7</td>
</tr>
<tr>
<td>6</td>
<td>Flowchart</td>
<td>10</td>
</tr>
</tbody>
</table>
1. **Purpose**

The purpose of this Standard Operating Procedure (SOP) is to describe action(s) to be taken by the IEC when investigator(s)/ trial site(s) fail(s) to:

- follow the procedures written in the approved protocol,
- comply with national and/or international guidelines, statutory provisions, institutional guidelines or rules or procedures mandated by the Institutional Ethics Committee (IEC) for the conduct of human research,
- respond to the IEC requests regarding statutory, ethical, scientific or administrative matters.

2. **Scope**

This SOP applies to all IEC approved research protocols involving human research participants.

3. **Responsibility**

The IEC Secretariat is responsible for receiving deviation/violation reports as per (AX 01/SOP 11/V1) submitted by the Principal Investigator (PI)/others and placing it on the agenda of the meeting. Reporting of deviation/violation in any other reporting format will not be accepted.

The IEC members should review and take action on such reports.

4. **Definitions**


**Protocol Deviation and Protocol Violation:**

**Protocol Deviation** - A protocol deviation is any change, divergence, or departure from the study design or procedures of a research protocol that is under the investigator’s control and that has not
been approved by the IRB. Upon discovery, the Principal Investigator is responsible for reporting protocol deviations to the IRB using the standard reporting form.

**Protocol Violation**- A protocol violation is a deviation from the IRB approved protocol that may affect the subject's rights, safety, or wellbeing and/or the completeness, accuracy and reliability of the study data. If the deviation meets any of the following criteria, it is considered a protocol violation. Example list is not exhaustive.

I. The deviation has harmed or posed a significant or substantive risk of harm to the research subject. For example
   - A research subject received the wrong treatment or incorrect dose.
   - A research subject met withdrawal criteria during the study but was not withdrawn.
   - A research subject received an excluded concomitant medication.

II. The deviation compromises the scientific integrity of the data collected for the study. For example
   - A research subject was enrolled but does not meet the protocol's eligibility criteria.
   - Failure to treat research subjects per protocol procedures that specifically relate to primary efficacy outcomes. (if it involves patient safety it meets the first category above)
   - Changing the protocol without prior IRB approval.
   - Inadvertent loss of samples or data.

III. The deviation is a willful or knowing breach of human subject protection regulations, policies, or procedures on the part of the investigator(s). For example
   - Failure to obtain informed consent prior to initiation of study-related procedures
   - Falsifying research or medical records.
   - Performing tests or procedures beyond the individual's professional scope or privilege status (credentialing)

IV. The deviation involves a serious or continuing noncompliance with federal, state, local or institutional human subject protection regulations, policies, or procedures. For example
   - Working under an expired professional license or certification
   - Failure to follow federal and/or local regulations, and intramural research or CC policies
   - Repeated minor deviations.
V. The deviation is inconsistent with the NIH Human Research Protection Program’s research, medical, and ethical principles. For example
   • A breach of confidentiality.
   • Inadequate or improper informed consent procedure.

**Minor Protocol Deviation**- A minor protocol deviation is any change, divergence, or departure from the study design or procedures of a research protocol that has not been approved by the IRB and which DOES NOT have a major impact on the subject's rights, safety or well-being, or the completeness, accuracy and reliability of the study data.

5. Detailed instructions

5.1 Detection of Protocol deviation/ violation

Protocol deviation/ violation may be detected in one the following ways (but not limited to those listed below):

a. Protocol deviation/ violation may be reported by Investigator/ study site/ sponsor/ Contract-Research Organization to the IEC.

b. The IEC members performing monitoring of the project at trial site may detect protocol deviation/violation if the project is not been conducted as per protocol/ national/ international regulations.

c. The Secretariat may detect protocol deviation/ violation from failure to comply with statutory requirements/ failure to respond to requests from IEC within reasonable time limit/ failure to respond to communication made by IEC.

d. The IEC members may detect protocol deviation/ violation when scrutinizing annual/ periodic reports/ SAE reports/ any other communication received from the Investigator/ trial site/ sponsor/ study monitor/ contract research organization.
e. The IEC Secretariat and/or IEC members may become aware of a protocol deviation/violation while reviewing study-related documents including reports filed in by the Principal Investigator (PI).

f. Communication/complaint/information received from a research participant who has been enrolled or any individual who has been approached for enrollment.

g. Any report/communication brought to the notice of Member, Secretary/Jt. Secretary/Chairperson of IEC by an independent person.

h. Communication received from the Head of the Institution informing IEC about an alleged protocol violation/protocol deviation.

5.2 Receipt of protocol deviation/violation report by the Secretariat

1. The PI will report the protocol deviation/violation as per Annexure 1 AX 01/SOP 11/V1.

2. In case protocol deviation/violation is detected by any other person (See Section 5.1) and reported to the IEC (there is no format for this), the Member Secretary will write to the PI to submit a protocol deviation/violation as per Annexure 1 AX 01/SOP 11/V1.

3. The Secretariat will notify the Member Secretary of any protocol deviation/violation report received from the PI/from any source within 2 working days of receipt of the notification.

5.3 Actions to be taken

1. The action of the IEC will be based on:
   o The nature and seriousness of the deviation/violation.
   o Frequency of deviation/violation in the study in the past.
   o Frequency of deviation/violation in previous studies conducted by the same PI/Co-PI or in the same department.

2. Member Secretary will decide on the impact of the protocol deviation/violation and act accordingly. Depending upon the seriousness, the IEC shall do the following (not limited to these actions):
o Ask PI for written clarification as soon as the deviation is received

o If the impact is serious, this report will be shared with the Chairperson and two or more IEC members designated by the Chairperson.

o If the impact of the protocol deviation is serious enough, the Member Secretary will instruct the Secretariat to call for and schedule a full-board meeting specifically to discuss the issue within 7 working days of the initial scrutiny

o The Secretariat will put up the information and communication at the next full board meeting for discussion.

3. The Member Secretary in consultation with IEC members will review the information available and deliberate on it.

4. The Chairperson will take a final decision depending on the seriousness of the violation. The decision will be taken to ensure that the safety and rights of the research participants are safeguarded. The decision will be taken by voting. A majority vote for approval, disapproval or request for modifications of a study suspension or termination of an ongoing study is defined as 2/3rd of the voting members present at the meeting.

5. The decision taken by IEC could include one or more of the following:

   o Determine that no further action is required, or take other actions as appropriate.

   o Inform the PI that the IEC has noted the violation / deviation, and instruct the PI to ensure that deviations/ violations do not occur in future and to follow IEC recommendations.

   o Enlist measures that the PI would undertake to ensure that such deviations / violations do not occur in future.

   o Observe the research or consent process (depending on the nature and frequency of the deviation).

   o Suggest modifications to the protocol.
FERCI MODEL SOPs

Review of Protocol Deviations / Violations

SOP 11/V1:

Effective Date:
aa/bb/cccc

- Alter the interval for submission of the continuing review/annual project status.
- Ask for additional training of the investigator and study team.
- Reprimand the PI.
- Seek additional information from the PI.
- Conduct audit of trial by the IEC.
- Suspend the study till additional information is made available and scrutinized.
- Suspend the study till recommendations made by the IEC are implemented by the PI and found to be satisfactory by the IEC.
- Suspend the study for a fixed duration of time.
- Suspension or termination of the study.
- Revoke approval of the current study.
- Inform DCGI/other relevant regulatory authorities.
- Keep other research proposals from the PI/Co-PI under abeyance. Review and/or inspect other studies undertaken by PI/Co-PI.

6. This final decision will be recorded on AX 01/SOP 11/V1 by the Member Secretary.

5.4 Procedure for notifying the PI and other concerned authorities

- The Member Secretary will draft a notification letter.
- The signed letter by Member Secretary will be sent to the PI and Department Head(s) (if required on case to case basis) and Institutional Officials (if required on case to case basis).
- The IEC secretariat will send a copy of the notification to the relevant national authorities (if required on case to case basis) and institutes (if required on case to case basis in case of multi-centric trials).

5.5 Records and follow up to be kept by IEC secretariat

- The Secretariat will keep a copy of the notification letter in the respective project file.
6. Annexure

Annexure 1  AX 01/SOP 11/V1 - Deviation/ Violation Record

<table>
<thead>
<tr>
<th>IEC Protocol no.:</th>
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<tbody>
<tr>
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<table>
<thead>
<tr>
<th>Study Title:</th>
</tr>
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<tbody>
<tr>
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<table>
<thead>
<tr>
<th>Principal Investigator:</th>
</tr>
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<tbody>
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<td></td>
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<table>
<thead>
<tr>
<th>Department:</th>
</tr>
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<tbody>
<tr>
<td></td>
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- ☐ Deviation from protocol
- ☐ Violation

<table>
<thead>
<tr>
<th>Description of deviation (s)/violation(s)</th>
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<tbody>
<tr>
<td></td>
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Corrective Actions Taken by the Principal Investigator:

- __________________________________________
- __________________________________________
- __________________________________________

Reported by (Name of Principal Investigator/ Study Team Member):

- __________________________________________

Signature with date: _________________________
**Provisional Decision by the Reviewer (Member Secretary and/or Chairperson and/or IEC Member/s)**

- [ ] Noted
- [ ] Request the PI not to perform such deviations/ non compliances/ violations in future
- [ ] Specific recommendations stated below to be followed

<table>
<thead>
<tr>
<th>Reason for termination</th>
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<tbody>
<tr>
<td>Suspend the study till the IEC recommendations are implemented</td>
</tr>
<tr>
<td>Suspend the study till information available</td>
</tr>
<tr>
<td>Terminate approval of the current study</td>
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**Reviewed by**

Name/s:

Signature/s with date:
Discussion of the protocol deviation/violation at the

☐ Emergency meeting on _________________

☐ Next Scheduled full board meeting on _________________

Final decision at the full board meeting held on

___________________________________________

Signature with date
IEC Member Secretary

7. Flow Chart

<table>
<thead>
<tr>
<th>No.</th>
<th>Activity</th>
<th>Responsibility</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Detection and reporting of Protocol deviation/</td>
<td>IEC members/ Secretariat/ principal investigator</td>
</tr>
<tr>
<td></td>
<td>violation</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Receipt of protocol deviation / violation report</td>
<td>Secretariat</td>
</tr>
<tr>
<td>3</td>
<td>Review, board discussion, decision and action</td>
<td>IEC Members, Member Secretary and Chairperson</td>
</tr>
<tr>
<td>4</td>
<td>Notify the Principal Investigator/ concerned</td>
<td>Secretariat</td>
</tr>
<tr>
<td></td>
<td>authorities of IEC action</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Maintain records</td>
<td>Secretariat</td>
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</table>