Title: Continuing Review of Study Protocols

SOP Code: SOP 10/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]

Prepared by:

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<tr>
<th>Name</th>
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<tr>
<td>Dr. Padmaja Marathe, FERCI Member</td>
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Reviewed by:

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<tr>
<td>Dr. U. M. Thatte, FERCI Secretary</td>
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<td>Dr. S. K. Kamat, FERCI Treasurer</td>
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Approved by:

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<tr>
<td>Dr. Vasantha Muthuswamy, FERCI President</td>
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1. Purpose

The purpose of this Standard Operating Procedure (SOP) is to describe how continuing review of previously approved protocols should be managed by the Institutional Ethics Committee (IEC). The purpose of the continuing review is to periodically monitor the progress of the study, to ensure continuous protection of the rights and welfare of research participants.

2. Scope

This SOP applies to conducting any continuing review of already approved study protocols at pre-specified intervals. All the projects approved by the IEC will be reviewed at least once a year. Depending upon the degree of risk to the participants, the nature of the studies, the vulnerability of the study participants and duration of the study, the IEC may choose to review or monitor the protocols more frequently.

3. Responsibility

It is the responsibility of the IEC Secretariat to remind the PIs and Member Secretary regarding continued review of protocols at the correct interval. All the approved protocols will be reviewed annually. It is the responsibility of the Member Secretary to ensure a decision regarding whether the project needs to be reviewed more frequently is taken during the IEC meeting in which the project is finally approved. This must be recorded in the minutes. A fresh decision to increase review may be taken if required based on the SAE reports, monitoring reports, or safety concerns. This is responsibility of the SAE subcommittee and Member Secretary.

The IEC is responsible for reviewing the progress made in the protocol (number of patients recruited, dropped out, reasons for drop-out), the occurrence of unexpected events or problems, and compliance of the investigator regarding IEC communication.
4. Detailed instructions

4.1 Determining the date of continuing review

- The date of the continuing review will always be at least once in the year.
- The IEC may recommend more reviews during the approval process depending on the level of risk. This will be documented in the minutes.
- The Secretariat will inspect the minutes of meeting to set a timetable for continuing review.
- The Secretariat will identify and record the due dates for each project.

4.2 Notifying the PI or the study team

The Secretariat will send a reminder to the PI as per the format AX 01/SOP 10/V1 two months prior (if an annual review) or less as appropriate (if any special additional reviews are required) to the due date of continuing review.

4.3 Managing the continuing review package upon receipt

- The Secretariat will receive a package (soft and hard copy) submitted by the PI for continuing review of each approved protocol. Only one set (soft and hard copy) of continuing review report shall be submitted by the PI to the IEC as per the format Continuing Review Application Form (AX 02/SOP 10/V1).

4.4 Verifying the contents of the package

- The Secretariat will ensure that the contents of the package include the following documents:
  - Continuing Review Application Form (AX 02/SOP 10/V1)
  - The Continuing Review Application Form duly filled with an explanation for any “yes” (ticked on the Continuing Review Application Form (AX 02/SOP 10/V1))
answers on the application form and a discussion of scientific developments, either through the conduct of this study or similar research that may alter risks to research participants. The changes in the selection criteria of participants, protocol/Informed consent Document amendments, changes in the study team, any unexpected complications etc. must have been discussed in the attached narrative.

- The Secretariat will confirm complete information is appended and for the presence of the required signatures of the Principal Investigator in the Continuing Review Application Form (AX 02/SOP 10/V1).

### 4.5 Review process

- The Continuing review submission may undergo expedited review (as per the procedure described in SOP7B/V1) or full board review (as per the procedure described in SOP7A/V1) as deemed appropriate by the IEC Chairperson/ Member Secretary
- The IEC Chairperson/ Member Secretary/ Member/s will use the Continuing Review Application Form (AX 02/SOP 10/V1) to guide the review and deliberation process.
- The Secretariat will send the Continuing Review Application Form (AX 02/SOP 10/V1) to the designated IEC members.
- The IEC Chairperson/ Member Secretary/ Member/s could reach one of the following decisions after review:
  1. Noted - The IEC approves the continuation of the project without any modifications.
  2. Modifications recommended: The study protocols that have been suggested modifications by the IEC may not proceed until the conditions set by the IEC in the decision have been met. The amendments and the required documents should be amended and submitted to the IEC within one month for re-review.
3. The project cannot be continued: The reasons for discontinuation of the project should be mentioned in the letter notifying the decision to the Principal Investigator. This decision shall be recorded by the Member Secretary on AX 02/SOP 10/V1.

- The IEC Chairperson will sign and date the IEC decision on Continuing Review Report after a decision has been reached.
- The decision on continuing review taken by the Chairperson/ Member Secretary/ Member/s will be informed to all IEC members at the next full board meeting.
- The continuing review report may be discussed at full board if deemed necessary by Chairperson/Member Secretary.
- The IEC Secretariat will maintain and keep the IEC Decision forms and minutes of the meeting relevant to the continuing review as part of the official record of the review process in the project file.

4.6 Communicating IEC Decision to the PI

- The Secretariat will notify the PI of the decision within 14 days of the meeting at which the report was discussed or of the date of review by the Chairperson/ Member Secretary/ IEC Member/s.

4.7 Non-submission of continuing review report by principal investigator before due date.

- If a PI fails to submit the continuing review report within one month of the due date (i.e. 11th months from the date of approval, or earlier on the dates as specified), the Secretariat will send a telephonic and/or email reminder at least 15 days prior to due date of review. If there is no response, the IEC secretariat will put up the matter for discussion at the forthcoming full board meeting for appropriate action which may consist of but not limited to sending:
  a) A reminder letter again
  b) A letter asking explanation for non-submission
c) A letter asking the PI to put recruitment of new participants on hold till report is submitted

d) Any other action as deemed appropriate by IEC

5. Annexures
Annexure 1  AX 01/SOP 10/V1- Reminder letter by the IEC to principal investigator
Annexure 2  AX 02/SOP 10/V1- Continuing Review Application Form

Annexure 1: AX 01/SOP 10/V1

Reminder letter by the IEC to principal investigator

Date:-
Name of Principal Investigator:-
Department:-
Ref: - Project no.  Title: XXXXXX

The above referenced project was approved by the IEC on XXXXXXX and is due for Continuing Annual/ Periodic Review by the IEC. You are requested to submit an Annual/ Periodic status report in the prescribed format which is enclosed (Continuing Review Application Form AX 02/SOP 07) at the earliest, on or before XXXXX. (1 month period)

Signature with date ________________________________
Member Secretary/ Chairperson ________________________________

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<th>Project No.:</th>
<th>Date of IEC approval:</th>
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<tr>
<th>Principal Investigator :</th>
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Annexure 2: AX 02/SOP 10/V1

Continuing Review Application Form
## FERCI MODEL SOPs
### Continuing Review of Study Protocols

**SOP 10/V1**

**Effective Date:** aa/bb/cccc

### Summary of protocol participants:
- No. of participants screened
- No. of participants approved by IEC
- No. of recruited participants
- No. of ongoing participants
- No. of completed participants
- No. of participants who refused to consent

### Have any participants been withdrawn from this study?
- Yes [ ] No [ ]

If no, state the number and reasons for drop-outs of each participant, attach separate sheet if needed.

### Have there been any amendments in protocol/Informed Consent Document since the last review?
- Yes [ ] No [ ]

Were these protocol/Informed Consent Document (ICD) amendments approved by IEC?
- Yes [ ] No [ ]

If no, mention the amendments not approved.

Which protocol amendment is the site following at present ________________

Which ICD amendment is the site following at present ________________

### Has any information appeared in the literature, or evolved from this or similar research that might affect the IEC/IEC’s evaluation of the risk/benefit analysis of participants involved in this protocol?
- Yes [ ] No [ ]

If Yes (attach separate sheet if needed)

### Whether reports of SAEs so far have been reviewed by the IEC?
- Yes [ ] No [ ]

☐ Whether reports of SAEs at other sites have been submitted to the IEC-__________

### Have any participating investigators been added or withdrawn since last review?
- Yes [ ] No [ ]

If Yes (Identify all changes in the attached narrative)

### Is report of interim data analysis available?
- Yes (submit as an attachment) [ ] No [ ]

### Is report of the data safety and monitoring board available?
- Yes (submit as an attachment) [ ] No [ ]

### Have any investigators developed equity or consultative relationship with a source related to this protocol which might be considered a conflict of interest?
- Yes (Append a statement of disclosure) [ ] No [ ]

### Date: ________________

**Signature of the Principal Investigator with Date:** ________________

**Assessment of Continuing Review Report by the IEC**

To be reviewed by
FERCI Model SOPs

Continuing Review of Study Protocols

SOP 10/V1
Effective Date: aa/bb/cccc

- Chairperson /Member Secretary only and informed to the IEC members at Full Board
- Full Board
- Any 2 IEC members and informed to the IEC members at Full Board
  1. Names of IEC members: _________________________________
  2. _________________________________

Signature with date
Chairperson/ Member Secretary

IEC Decision on the Continue Review Report

Date - ------------------

Decision
Approved and the project can be continued without any modifications
Modifications recommended - requiring protocol resubmission
State the recommendations:

Protocol should be discontinued
State the reasons for discontinuation

Date of Full Board discussion

Signature of reviewer/s with date: _________________________________

Signature with date
Chairperson / Member Secretary
6. Flow Chart

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<tr>
<th>No.</th>
<th>Activity</th>
<th>Responsibility</th>
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<tbody>
<tr>
<td>1</td>
<td>Determine the date of continuing review</td>
<td>Administrative Officer / Secretariat</td>
</tr>
<tr>
<td>2</td>
<td>Notify the Principal Investigator or study team</td>
<td>IEC Secretariat</td>
</tr>
<tr>
<td>3</td>
<td>Manage continuing review package upon receipt and verifying its contents</td>
<td>IEC Secretariat</td>
</tr>
<tr>
<td>4</td>
<td>Notify the members of the IEC</td>
<td>IEC Secretariat</td>
</tr>
<tr>
<td>5</td>
<td>Review of Continuing review report</td>
<td>IEC Secretariat, Members and Chairperson</td>
</tr>
<tr>
<td>6</td>
<td>Prepare meeting agenda</td>
<td>IEC Secretariat</td>
</tr>
<tr>
<td>7</td>
<td>Communicate the IEC decision to the Principal Investigator</td>
<td>IEC Secretariat</td>
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