

SOP Code: SOP 13/V2 Effective from 21/10/2019

Title: Review of Study Completion Reports

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Effective Date: 21-10-2019

SOP Constitution and Approval:

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1. Purpose

The purpose of this Standard Operating Procedure (SOP) is to provide instructions on the review of Study Completion Report submitted for studies approved by the Institutional Human Ethics Committee (IHEC), MGMCRI.

2. Scope

This SOP applies to the review of Study Completion Report which is a written report of every completed study submitted by Principal Investigator (PI).

3. Responsibility

It is the responsibility of the Secretariat/ IHEC Chairperson/ Member Secretary/Additional Member Secretary/ Member/s to review the study report and act on it.

4. Detailed instructions

4.1. Receipt of Study Completion Report

- The Secretariat will receive 1 copy (soft and hard) of Study Completion Report filled as per the format AX 01/SOP 13/V2 from the PI. The study completion report is expected from the investigator within 1 month of completion of the study at the site.
- The Secretariat will follow instructions as in "Management of Protocol Submission" for receiving and checking the report package.
- It is the responsibility of the IHEC Secretariat to review the report for completeness.
- The Secretariat shall verify the submitted Study Completion Report along with Study Completion Report Form- AX 01/SOP 13/V2 and forward it to the Member Secretary/ Additional Member Secretary within 7 working days of receipt.
- The Member Secretary/ Additional Member Secretary will review the Study Completion Report, confirm that it is complete and present it at the next full board meeting.
- If there is a need felt (e.g. a deviation/ violation is noted), the Member Secretary/ Additional Member Secretary will handle it as per SOP 11/V2.



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 The Secretariat shall include the Study Completion Report Form in the agenda for IHEC members for discussion at the full board meeting.

4.2. During the Board meeting

- The Member Secretary/ Additional Member Secretary will present the report and members can discuss as needed.
- Following the discussion, the Chairperson may take one of the following decisions:
 - a) noted / approved
 - b) request for additional information / clarification
- The Secretariat will note the decision in the meeting minutes.
- The Member Secretary/ Additional Member Secretary will draft a letter to the PI conveying decision on the study completion report.
- The study shall be considered as closed if the decision by IHEC is "Noted" or "Approved".
- The Secretariat will accept and file the report and get the Study Completion Report Form AX 01/SOP 13/V2 signed by the Chairperson.
- The final report will be placed in the master file and kept in the archival area.
- The Administrative Officer will archive the entire study for a period of 5 years from the date of completion of the project if the decision is noted and closed.

5. Reference to other applicable SOPs:

SOP 06/V2: Management of Research Study Protocol and study Related Documents Submitted for Ethics Review.

SOP 08/V2: Agenda Preparation, Meeting Procedures and Recording of Minutes

SOP 11/V2: Review of Protocol Deviations / Violations

6. Annexures

Annexure 1: AX 01/SOP 13/V2 - Study Completion Report



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Annexure 1: AX 01/SOP 13/V2

Study Completion Report

M	GMCRI-IHEC Ref. No. (for office use):	
Ti	tle of study:	
Pr	incipal Investigator (Name, Designation and Affiliation):	
1.	Date of EC approval:	
2.	Date of start of study: Date of study completion:	
3.	Provide details of:	
	a. Total number of study participants approved by the EC for recruitment:	
	b. Total number of study participants recruited:	
	c. Total number of participants withdrawn from the study (if any):	
Pr	ovide the reasons for withdrawal of participants (Explanation for the withdrawal of participants	
wh	ether by self or by the PI):	
4.	Describe in brief the publication/ presentation/dissemination plans of the study findings.	
	(Also, mention if both positive and negative results will be shared)	
5.	Describe the main ethical issues encountered in the study (if any)	
6.	State the number (if any) of Deviations/Violations/ Amendments made to the study protocol during the study period	
Dε	eviations: Violation: Amendments:	
7.	Describe in brief plans for archival of records / record retention:	
8.	Is there a plan for post study follow-up? Yes No	
If	ves describe in brief	



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9. Do you have plans for ensuring that the data from the study can be shared/ ac	cessed easily?
If yes, describe in brief:	Yes No
10. Is there a plan for post study benefit sharing with the study participants? No	☐ Yes☐
If yes, describe in brief:	
11. Describe results (summary) with Conclusion (For sponsored studies, if the final rep from sponsor, it may be submitted later to the EC once it is ready):	ort is not available
12. Number of SAEs that occurred in the study:	
13. Have all SAEs been intimated to the EC? No	Yes Yes
14. Is medical management or compensation for SAE provided to the participants No	s? Yes
If yes, provide details	
Signature of PI with date:	



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7. Flow chart

Receipt of the study completion report IHEC Secretariat

Checking the contents of the report packages and assess adequacy of contents IHEC Secretariat

Verification of the study completion report, preparation of the study completion statement and sending them to the Member Secretary

IHEC Secretariat

Review of Study completion report for completeness and informing members at full-board meeting

Member-Secretary/Additional Member Secretary Chairperson

Inclusion of report/ review at full-board meeting IHEC Secretariat

Discussion and decision at the full board meeting
Member Secretary/Additional Member Secretary / Chairperson

Noting the decision in the minutes of the Meeting IHEC Secretariat

Conveying decision to the Principal Investigator IHEC Secretariat

Archiving all the study-related documents along with the Study completion report

Administrative Officer

8. References

- Forum for Ethics review Committees in India (FERCI). Standard Operating Procedures
 of Institutional Ethics Committee (cited 5th October 2019). Available from:
 http://www.ferci.org/sops/
- Indian Council of Medical Research (ICMR). National Ethical guidelines for biomedical and health research involving human participants, October 2017 (cited 6th October 2019) available from: http://www.icmr.nic.in/guidelines/ICMR Ethical Guidelines 2017.pdf