Title: Preparing for monitoring and audit

SOP No.: 19/05

Date first effective: 01 Jan 2023 Review date: 31 Dec 2023

Department of Clinical Pharmacology, 1st Floor, New MS Building, Seth GS Medical College & KEM Hospital, Parel, Mumbai 400012.

Category: Study conduct

Title: Preparing for Monitoring and audit

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Authors: Dr. Dhruve Soni

DM Resident

Signature with date

21/ DEC/ 2022

Reviewer:

Dr. Vijaya Gunjal Assistant Professor

Signature with date

(31/0Ec/2022

Approved by:

Dr.Nithya Gogtay Professor and Head

Signature with date

W 31. 12. 22

Dr. Nithya Gogtay Professor & Head

Department of Clinical Pharmacology

1" Floor, MS Building,

Seth GS Medical College & KEM Hospital

Parel, Mumbai - 400 012.

Department of Clinical Pharmacology

Seth GS Medical College & KEM Hospital,

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1. Purpose:

This standard operating procedure (SOP) describes the responsibilities of the PI and study team in its preparation for a monitoring and audit visit.

2. Scope:

This SOP is limited to the responsibilities of the study team in its preparation for a monitoring and audit visit.

3. Responsibilities:

Principal investigator, Co-investigator, Study Coordinator or any other appropriately qualified staff in the team, as delegated by the Principal Investigator, will be responsible for implementation of this SOP.

4. Applicable rules, regulations and guidelines

- Indian GCP Guidelines 2001
- Ethical Guidelines for Biomedical and Health Research involving Human Participants, ICMR 2017
- ICH E6(R3) EWG Draft Guidelines dated 19th April, 2021.

5. Refences to other SOPs:

SOP No. 18/05: Archiving

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6. Detailed instructions

- 1. The purpose of study monitoring/audit is to verify that:
- · The rights, safety and well-being of human participants are protected
- Reported data are accurate, complete and verifiable
- The study is conducted in compliance with the protocol and applicable guidelines and regulations.
- The PI must permit monitoring and audit by IEC members, regulators and/ or sponsors at all times that is asked by the authorities.
- 3. At the same time the PI and study team should ensure that no document (original or copy) which allows the identification of a participant in the study is shared with the inspectors unless insisted upon by regulators and IEC members. Sponsors representatives should at no time be given access documents that permit dentification of a participant.
- 4. The PI should be available throughout the monitoring.
- The study coordinator should ensure availability of a suitable location for monitoring.
- The PI and study coordinator must ensure the availability of all team members for the monitoring.
- The study coordinator must ensure that all relevant documents of the study (including but not limited to the following) are available for the monitor viz.
- Trial master file(s)
- Case Report Forms and source notes
- Participant Informed Consent Forms
- Participant's medical files
- Documents related to the Investigational product
- Documents related to the sample collection, storage and shipment
- All documentation related to AE/SAEs

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- Documentation or correspondence with Ethics Committee
- 8. Study coordinator should ensure that all documentation (including lab reports) is complete including PI signatures, prior to monitoring/audit. Signature of the PI should be obtained in real time.
- The study coordinator should confirm the identity of monitors upon arrival by checking identity cards.
- 10. Measures to ensure site preparedness for a monitoring visit or an audit are mentioned in Appendix 1 and 2.

7. Appendices

Appendix 1: Preparedness for monitoring by the sponsor [1]

	How to ensure site preparedness	
Pre-Study monitoring/ visit	 Keep the SOPs of the site ready and ensure that all SOPs are current and valid Keep the IEC SOPs / URL where they can be found ready Ensure that all instruments relevant to the study e.g. refrigerators, centrifuges ECG machine, weighing balance, BP apparatus, height measuremen apparatus [as applicable to the study] are calibrated and the calibration certificates are ready for inspection by the monitor Ensure documentation for controlled access (Audio-Video consenting area pharmacy room, document archival area, clinical pharmacology unifoutpatient department etc.) for all study areas is available for examination be monitor Ensure documentation for updated emergency tray and all requisit instrumentation calibration and functioning (for example ECG, defibrillato ventilator) so that these can be checked and verified by the monitor Ask for a report of the pre-study visit 	

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Site –	 Prepare specific questions related to operational aspects of the study which 	
nitiation	can be discussed with the sponsor at the time of the site initiation	
visit	 Ensure that the entire team is present on the day of the site initiation 	
	 Confirm supplies received [investigational product (IP), Trial master file 	
	(TMF), laboratory kits etc.] from the sponsor/ CRO	
	 Ensure that any deficiencies identified during the pre-study visit have been 	
	addressed	
	 Ask for a report of the site initiation visit report. When received, file in the 	
	Trial Master File	
Routine	Confirm a clear understanding among the team members of individual roles	
monitoring	and responsibilities	
visit	Ensure that all study team members are available	
	Ensure a quiet area is available for the monitoring	
	• Ensure proper documentation of Case Report Form (CRF) including	
	signatures, updated TMF and other study related logs	
	 Arrange all case sheets, CRFs and TMFs clearly and sequentially 	
	 Ensure that all documents are returned to their original place after monitoring 	
	Enquire regarding findings at the end of the visit	
	 Ensure that medical records and files are kept in a locked room, if monitorin 	
	lasts for multiple days	
	 Discuss the findings in a formal interview and resolve as many findings a 	
	possible	
	Tentative dates for the next visit may be discussed	
	• File the monitoring report/ follow up reports sent by the monitor in the TM	

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Close out	Ensure that all study team members are available
visit	Make necessary arrangements for retrieval of study related documents
	• Provide secure area for archiving documents for a specific period as per
	sponsor's SOP
	 In the case of hard copies, confirm that all case report forms are retrieved and
	submitted to sponsor and copies archived.
	 Ensure that all extra CRFs, study supplies and laboratory kits returned to the
	sponsor/CRO
	 Ensure that all biological samples have been shipped or back-up samples are
	destroyed as per site SOP and protocol
	 Make sure that the final report provided by the monitor is placed in the TMF.
	All the electronic data should be archived as mentioned in the protocol and
	as per sponsor policy
	Send a copy of the final monitoring report to the IEC and close the study with
	EC
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Appendix2: Site Preparedness for a sponsor or a regulatory audit

How to ensure preparedness

- Ensure a secure room and study team availability on the day of audit
- Ensure that the TMF is updated per site SOP
- Ensure subject screening /enrollment log and the duty delegation log are up to date
- Keep the initial IEC approval letter in the TMF and latest amendment approvals if changes have been made to the study

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- Ensure that all correspondence (signed/dated applications, responses, e-mails)
 to and from the IEC and sponsor are filed
- Ensure re-consenting [if applicable] has been completed and documented
- Make sure protocol deviation /violation report have been submitted to the IEC and the IEC correspondence is filed in the TMF
- Ensure all IEC correspondence of SAE report(s), if any, are available in the TMF
- Ensure data collection, source documents and IP accountability log for each participant are up to date
- Ensure that samples collection, storage and shipment logs are updated
- Make sure that the audit log is up to date in case the site was audited previously
- Keep all study hard copies in a cupboard with restricted access
- Ensure that access to electronic study records and files are password protected

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8. Glossary

Definition of Monitoring:

The act of overseeing the progress of a clinical trial, and of ensuring that it is conducted, recorded, and reported in accordance with the protocol, standard operating procedures (SOPs), GCP, and the applicable regulatory requirement(s). ICH E6(R3) EWG Draft Guidelines dated 19th April, 2021.

Definition of Audit:

An audit is a systematic and independent examination of trial related activities and documents to determine whether the evaluated trial related activities were conducted, and the data were recorded, analyzed and accurately reported according to the protocol, sponsor's SOPs, GCP, and the applicable regulatory requirement(s). ICH E6(R3) EWG Draft Guidelines dated 19th April, 2021.

Reviewer:

Dr. Vijaya Gunjal Assistant Professor

Signature with date

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Department of Clinical Pharmacology Seth GS Medical College & Hos Parel, Mumbai - 45 912.

Approved by:

Dr.Nithya Gogtay Professor and Head

Signature with date 31.12.42

Dr. Nithya Gogtay Professor & Head Department of Clinical Pharmacology 1" Floor, MS Building, Seth GS Medical College & KEM Hospital Parel, Mumbai - 400 012.