Standard Operating Procedures for Institutional Ethics Committee

Rajendra Institute of Medical Sciences, Ranchi

Standard Operating Procedures for Institutional Ethics Committee, Rajendra Institute of Medical Sciences, Ranchi

I. Standard Operating Procedure (SOP)

IEC, RIMS, RANCHI

Version: 4.0

Date: 15.09.2021

Pages: 32, Annexures: 19

II. SOP prepared by:

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III. SOP reviewed and approved by Institute Ethics Committee

Name and Designation	Signature with date
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IV. SOP accepted by:

Name and Designation	Signature with date
Dr. (Prof) Kameshwar Prasad Director RIMS, Ranchi	15/09/2001

Serial No.	Current Version Number	Effective Date	Description (Changes from the previous)
1	1.0	22 nd January 2016	Not Applicable
2	2.0	20 th September 2018	Change in SOP Sl. No. 13 Recordkeeping and archival, page No. 14, point No. 3 is changed from 3 years to 5 years
3	3.0	25 th December 2020	 i. Introduction of "Preparing SOPs" page No. 4-8 ii. Introduction of "General Principles for functioning" Page No.9-10 iii. Introduction of "Informed Consent" Page No. 11-12 iv. Introduction of "Special considerations/Protection of Vulnerable Population" Page No. 13-16 v. Change in Composition of IEC, RIMS, Ranchi Page 18 vi. Introduction of "Office bearers' and Member Specific Roles and Responsibilities" Page No. 19-23 vii. Change in EC Review Fees, Amendment Fees and Archival Fee, Revised Initial review fee. Page No.23-25 viii. Introduction of "Review of Serious Adverse Event Reports and Compensation issues" Page No. 27-29 ix. Introduction of "Review of the Study Completion Report" Page No. 29 x. Introduction of "Waiver of the Informed Consent" Page No. 31 xi. Accountability of researchers for the protection of the environment and resources. Point 6; Page No. 32 xii. Introduction of "Annexures 1 to 14" Page No 35-53
4.	4.0	15 th September, 2021	 i. References of Schedule Y replaced with New Drugs and Clinical trial rules, 2019. ii. Informed Consent (at number 5) and "Special considerations/Protection of Vulnerable Population" brought down (to page number 33-36) near waiver of consent;

Proforma for the Subject Information Sheet attached as Annexure 14

[Participant Informed Consent Form for participants more than 18 years of age (FORM 3A)] and Parents/Legally accepted representative (LAR) Consent *Form* (FORM 3B), Assent Form (FORM 3C) attached as annexure 15, 16 and annexure 17.

- iii. Office bearers' and Member Specific Roles and Responsibilities
- iv. Record Keeping and Archiving name changed to Maintenance of records by the IEC for clinical trials and points added.
- v. IEC approval notice for the studies in annexure 07 changed to the Format to accord approval to clinical trial protocol by the IEC, RIMS, Ranchi.
- vi. Recommendations of payments of compensation and determination The Quantum of Compensation in Cases of Clinical Trial related Injury or Death added in section 1
- vii. In section 13, Recommendations of payments of compensation in case of serious adverse event (SAE) criteria for assessment of trial relatedness of a serious adverse event, and determination the quantum of compensation in cases of Clinical Trial related Injury or death are added.
- viii. Form 2 (Form to be filled up by PI for submission to the IEC) added as annexure 10.
- ix. Checklist for verification of proposals submitted to IEC attached as annexure-19 (FORM 4).
- x. Record Keeping and Archiving at the office of IEC, RIMS, Ranchi named as Maintenance of records by Institutional Ethics Committee, RIMS, Ranchi for clinical trial (Section 20, page no. 40).

Table of Contents

1.	Preparing SOPs (Writing, Reviewing, Distributing & Amending SOPs) for the Institutional Ethics Committee (IEC), RIMS, Ranchi
2.	Objective
3.	General Principles for functioning
4.	Role and Responsibilities of IEC, RIMS, Ranchi
5.	Authority for constituting the IEC, RIMS, Ranchi
	Composition of IEC, RIMS, Ranchi
8.	Office bearers' and Member Specific Roles and Responsibilities
9.	Quorum Requirement
10.	Honorarium, Fees and Office expenses
11.	Application Procedure
12.	Review Procedure for research proposals

13. Review of Serious Adverse Event Reports and Compensation issues
13.1 Review of the SAE 13.2 Criteria for assessment of trial relatedness of injury or death or permanent disability to be related to clinical trial or bioavailability and bioequivalence stud 13.3 Recommendations of payments of compensation 13.4 Determination of the quantum of compensation in Cases of Clinical Trial related Injury or Death
14. Review of the Study Completion Report
15. Decision making and the format to accord approval to clinical trial protocol by the IEC, RIMS, Ranchi
16. Informed Consent
17. Special considerations/Protection of Vulnerable Population
18. Waiver of the Informed Consent
19. Responsibilities of Sponsors and Investigators towards IEC
20. Maintenance of records by Institutional Ethics Committee, RIMS, Ranchi for clinical trial
21. References
22. Annexures (1 to 19)
Annexure-1: Format for Appointment order for the members of IEC, RIMS, Ranchi.
Annexure-2: Terms of reference for the members, IEC, RIMS, Ranchi.
Annexure-3: Format for Consent Letter consent to be a member of IEC, RIMS, Ranchi.
Annexure-4: Confidentiality agreement for the members of IEC.
Annexure-5: Conflict of Interest (COI) Declaration form for the members of IEC.
Annexure-6: Format for the Office Order for the constitution of IEC, RIMS, Ranchi.
Annexure-7: Proforma to be submitted for MD/MS/DM/M.Ch/PhD/MBBS students' projects approval from IEC, RIMS, Ranchi

Annexure-8: Proforma for the submission of research proposals involving human participants for ethical approval from IEC, RIMS, Ranchi.

Annexure-9: Checklist for submission of research proposal to the IEC, RIMS, Ranchi.

Annexure-10: Form 2- Form to be filled up by PI for submission of research proposals to IEC, RIMS, Ranchi.

Annexure-11: Format to accord approval to Clinical Trial Protocol by the IEC, RIMS, Ranchi along with the undertaking by the investigator.

Annexure-12: Data Elements for reporting Serious Adverse Event occurring in a Clinical Trial/Bioavailability study/Bioequivalence study.

Annexure-13: Format for six monthly progress report of the Project.

Annexure-14: Proforma of the Subject Information Sheet

Annexure- 15: Consent Form for participants more than 18 years of age (FORM 3A)

Annexure-16: Consent Form for LAR for the children aged from birth to 16 years of age (FORM 3B)

Annexure-17: Assent Form for children aged 7 through 17 years of age and able to provide assent (FORM 2C)

Annexure-18: Informed document for Drug Clinical Trial [Checklist and Consent form] (FORM 2D, 2E)

Annexure-19: Checklist for the verification of proposals submitted to IEC, RIMS, Ranchi.

IEC, RIMS, RANCHI

1. PREPARING STANDARD OPERATING PROCEDURES (SOPs): WRITING, REVIEWING, DISTRIBUTING & AMENDING SOPS FOR THE INSTITUTIONAL ETHICS COMMITTEE (IEC), RIMS, RANCHI.

Purpose: This SOP defines the process for writing, reviewing, distributing, and amending SOPs within the IEC, RIMS, Ranchi.

The SOPs will provide clear, unambiguous instructions to conduct activities of the IEC in accordance with the ICMR guidelines 2006, WHO Operating Guidelines for Ethical Review Committee that Review Biomedical Research, and ICH (International Conference on Harmonization) Good Clinical Practice (GCP), Code Federal Regulations Title 21.

Scope: This SOP covers the procedures of writing, reviewing, distributing, and amending SOPs within the IEC, RIMS, Ranchi.

Roles and Responsibilities of the Members and the Secretariat: It is the responsibility of the chairperson of the IEC to appoint the SOP Team to formulate the SOPs. SOP team will prepare the draft SOPs. The draft SOPs will be reviewed and approved by the IEC members. The SOPs will then be signed by Director, RIMS, Ranchi. The SOP team will be responsible to amend the SOPs as and when required.

The SOP team will consist of Member Secretaries of IEC, administrative staff, and one or two other IEC members. The team will-

- Assess the request(s) for SOP revision in consultation with the Secretariat and Chairperson
- Propose a new, or modification in existing SOPs as needed
- Select the format and coding system for the SOPs
- Draft the SOP Review the draft SOP
- Submit the draft for approval to Chairperson.

Chairperson of the IEC:

Will appoint SOP Team
Will review and approve the SOPs
Will sign the approved SOP

IEC members:

Will review and sign the SOPs Will return all out of date SOPs to IEC office

Secretariat of IEC:

- Will co-ordinate activities of writing, reviewing, distributing, and amending SOPs.
- Maintain on file all current SOPs and the list of SOPs.
- Maintain a file of all SOP amendment requests
- Maintain an up-to-date distribution list of each SOP circulated to IEC members
- Maintain a record of the investigators to whom SOPs are distributed against a requisition if any
- Ensure that all IEC members and involved administrative staff have access to the SOPs
- Ensure that the IEC members and involved staff are working according to the current version of SOPs
- Maintain a file of all previous SOPs of the IEC
- Assist in the formulation of SOP procedure
- Ensure SOP revisions as and when required to comply with national regulations.

Detailed instructions

Identify the need for new or amendment to the SOP

Any member of the IEC, or administrative staff or investigators or administration can make a revision request or notices an inconsistency/ discrepancy / has any suggestions on how to improve the existing SOPs or requests to design an entirely new SOP, can put forth his / her request by using the Request Form for Formulation of new SOP/ Revision of an SOP Form. This Formulation of the new SOP/ Revision of an SOP Form is submitted to the Chairperson, IEC.

The Chairperson will inform all IEC members about this request in a regular full board meeting. If IEC members agree to the request, the Chairperson will appoint an appropriate SOP team comprising of Member Secretaries of both committees. The Chairperson may also appoint one or two committee members as members of the SOP team, if necessary. This designated team will proceed with the task of revision/formulation process of the SOP. If IEC members do not agree to the request, no further action will be taken. The Chairperson will inform the person/ IEC member who requested modification of the SOP in writing about the decision.

Appointment of the SOP team

The Chairperson will constitute an SOP team consisting of the Member-Secretaries administrative staff and one or two other IEC members who have a thorough understanding of the scientific and ethical review process. The SOP writing team will carry out the subsequent steps.

List of relevant SOPs

- Write down step by step all the procedures of the IEC
- Organize, devise and name each process

• Make a list of SOPs with coding format (ex- AX1-V3/SOP01/V3)

Design a format and layout

Each SOP should be given a number and a title that is self-explanatory and is easily understood A unique code number with the format SOP xx / Vy will be assigned to each SOP. xx is a two-digit number assigned to a specific SOP. "V" refers to the version of the SOP and "y" is a number identifying the version e.g.-SOP01/V4 is SOP number 01 with V=version no.04

Each Annexure (AX) is a unique code with format AXn–Vp/SOP xx/Vy. e.g. AX1–V4/SOP01/V4 indicates AX is Annexure, 4 is Annexure no., V4 is version 4, belonging to the SOP 01/V4 Each SOP will be prepared according to the template for Standard Operating Procedures (AX2 – V4/SOP01/V4). Each page of the SOP will bear a header with the effective date which is the date of approval of the SOPs by the Chairperson, IEC, and the Head of the Institution.

The SOP number will be on the left-hand corner of the header. The title of the SOP will be on the left-hand corner of the footer. The page number will be listed as Page—of---total pages on the right-hand- corner of the footer.

The first two pages of each SOP document will be signed and dated by the authors, the IEC members who have reviewed the SOPs, IEC Chairperson and Director, RIMS, Ranchi.

Preparation and submission of the final draft

- All the members of IEC may review the draft / revised SOP
- During respective IEC meetings, members can put forth their suggestions/comments on the draft / revised SOP
- The suggestions agreed upon unanimously by all IEC members will be incorporated and the final draft SOP will be formulated.
- The SOP team would stand automatically dissolved once the IEC takes the final decision regarding the SOP.

Final Approval of new/revised SOP

- The final version will be presented to the Chairperson of committee for review and approval. The Chairpersons will sign and date the SOP on the first page of the SOP document.
- This approved document will then be submitted to the Director, RIMS, Ranchi for acceptance. This date of approval is declared as the effective date for implementing the SOP.

Implementation, distribution, and filing of SOPs

- Approved SOPs will be implemented from the effective date.
- The Member Secretary will discuss the approved SOPs with the administrative staff and instruct them to implement the SOP accordingly.
- Approved SOPs will be distributed to IEC members and IEC staff according to the distribution list.

- When a revised version is distributed, the old version will no longer be effective. A copy of the old version will be archived in a master file.
- One complete original set of current SOPs will be archived in the SOP master file, by the IEC Secretariat and maintained in the IEC Office.
- A copy of the SOP master file will be maintained in the individual offices of IEC and DSMSC.
- Photocopies made from the paper versions of the SOP will be considered official only if stamped and signed by Member Secretary or authorized individual. A distribution log would be maintained.

Review and request for revision of an existing SOP

- Any member of the IEC or administrative staff or investigators or administration who notices
 that current SOPs have some lacunae or have any suggestions to improve a procedure should
 make a written request
- If IEC agrees with the request, the Chairperson will appoint an appropriate team for the revision process. If the committee does not agree, the Chairperson will inform the concerned individual who requested revision.
- The Member Secretary initializing the review and the Secretariat assists the Member Secretary of the SOP at least once every 2 years and records the dates of review in the SOP master file.

Manage and archive old SOPs

Old SOPs should be retained and marked "superseded" and archived in a file by the secretariat. The process of evolution of previous SOPs of the IEC will be documented in a defined format.

References

- 1. ICMR Ethical Guidelines for Biomedical Research on Human Participants, ICMR (2006)
- 2. WHO Operational Guidelines for Ethical Review Committee that Review Biomedical Research (Geneva 2000)
- 3. International Conference on Harmonization, Guidance on Good Clinical Practice (ICH GCP) (1996)
- 4. Code Federal Regulation Title 21
- 5. TMC IEC SOP 2016
- 6. AIIMS Raipur IEC SOP, 2021.
- 7. New Drugs and Clinical Trial Rules, 2019, CDSCO.

Standard Operating Procedures for the Institutional ethics committee (IEC) of RIMS, Ranchi

Rajendra Institute of Medical Sciences, Ranchi hereinafter referred to as "RIMS, Ranchi" has adopted these written Standard Operating Procedures (SOP) to ensure the protection of the rights and welfare of human participants in biomedical, experimental, and behavioral research conducted at RIMS, Ranchi.

2. OBJECTIVE

The objective of SOP is to ensure quality and consistency in ethical review of Biomedical Research Proposal in accordance with ICMR Ethical guidelines for biomedical research on human subjects and drugs and cosmetics act rules, Govt. of India.

3. GENERAL PRINCIPLES FOR FUNCTIONING OF IEC, RIMS, RANCHI:

Principle of essentiality: IEC, RIMS, Ranchi will consider the necessity of the use of human participants for the research.

Principle of voluntariness: IEC will ensure that the rights of the participants are safeguarded, informed consent is taken from all the participants in local language and that respect is given to participants' willingness or non-willingness to participate in the study.

Principle of non-exploitation: IEC will ensure that there is an equitable selection of the participants and the benefits and burdens of the research are distributed fairly. Sufficient safeguards to protect the **vulnerable groups** would be ensured. The vulnerable populations include children, pregnant and lactating women, people with racial inequalities, economically or socially disadvantaged people, mentally challenged and mentally differently-abled persons, and persons with reduced autonomy (prisoners, students, subordinates, employees, defense service personnel).

Principle of social responsibility: The IEC will ensure that the research is being conducted in such a way that in any way social harmony in community relationships is not disturbed.

Principle of ensuring privacy and confidentiality: IEC will ensure the privacy of the potential participants. Their identity and records would be kept confidential by the researcher and access will be limited to only those authorized. In some special circumstances for a valid scientific or legal reason, IEC will have the right to breach the privacy of the information.

Principle of risk minimization: Due care will be taken by all stakeholders (including researchers, ECs, sponsors, and regulators) of RIMS, Ranchi at all stages of the research to ensure that the risks are minimal and appropriate care and compensation is given if any harm occurs.

Principle of professional competence: IEC will ensure that the research is planned, conducted, evaluated, and monitored throughout by competent persons with appropriate and relevant qualifications, experience, and/or training.

Principle of maximization of benefit: IEC will ensure that the research is designed and conducted in such a way that the benefits to the research participants and the society are maximized.

Principle of transparency and accountability: The research plans and outcomes emanating from the research being carried out at RIMS, Ranchi would be brought into the public domain through registries, reports, and scientific and other publications while safeguarding the right to privacy of the participants. Stakeholders involved in research would disclose any existing conflict of interest and manage it appropriately. The research would be conducted in a fair, honest, impartial, and transparent manner to guarantee accountability. Related records, data, and notes would be retained for the required period for possible external scrutiny/ audit.

Principle of environmental protection: IEC will monitor (it may appoint a sub-committee to) monitor the researchers who will be accountable for ensuring the protection of the environment and resources at all stages of the research, in compliance with existing guidelines and regulations.

4. ROLES AND RESPONSIBILITIES OF IEC, RIMS, RANCHI

- **A.** The **IEC**, **RIMS**, **Ranchi** will review all types of research proposals involving human participants to safeguard the dignity, rights, safety, and wellbeing of all actual and potential research participants before approving the research proposals. The goals of research, however important, will never be permitted to override the health and wellbeing of the human participants.
- **B.** The **IEC**, **RIMS**, **Ranchi** will ascertain whether all the cardinal principles of research ethics viz., **autonomy**, **beneficence**, **non maleficence**, **respect for free and informed consent**, **respect for human dignity**, **respect for vulnerable persons**, **respect for privacy and confidentiality** as well as justice are taken care of in planning, conducting and reporting of the research.
- C. The IEC, RIMS, Ranchi will look into the aspects of protocol review, selection of participants, voluntary participation of potential participants, informed consent process, risk-benefit ratio, distribution of burden and benefit, maintenance of privacy and confidentiality, and provisions for appropriate compensation. It will review the proposals before the commencement of the study as well as during the study period through appropriate, well-documented procedures. The review will be based on the periodic study progress reports furnished by the investigators and/or monitoring and internal audit reports furnished by the sponsor and/or by visiting the study sites.
- **D.** It will review the proposals before the commencement of the study as well as review periodically until the completion of the study through appropriate, well documented procedures. such a review may be based on the periodic study progress reports furnished

by the investigators and/or monitoring and internal audit reports furnished by the sponsor and/or by visiting the study sites.

- **E.** The mandate of the **IEC** shall be to review all research projects to be conducted at the institution involving human beings directly or indirectly, irrespective of whether the research project is funded or non-funded, and if funded, then irrespective of the funding agency.
- **F. IEC, RIMS, RANCHI** will provide advice to the researchers on all aspects of the welfare and safety of the research participants after ensuring the scientific soundness of the proposed research.
- **G.** In case of **IEC**, **RIMS**, **RANCHI** revokes its approval according to a trial protocol, it will record the reasons for doing so and at once communicate such a decision to the investigator as well as to the licensing authority
- **H.** In case of a **serious adverse event** or death occurring to the clinical trial participant, **IEC**, **RIMS**, **RANCHI** shall forward it's reporting on the serious adverse event or death, after due analysis, along with its opinion on the financial compensation, if any, to be paid by the sponsor or his representative, whosoever had obtained permission from the licensing authority as defined under rule 21(b) for conducting the clinical trial, to the chairman of the expert committee constituted by the licensing authority under appendix xii (gazette notification 30th January 2013) with a copy of the report to the licensing authority within twenty-one calendar days of the occurrence of the serious adverse event of death.

5. AUTHORITY FOR CONSTITUTING THE IEC, RIMS, RANCHI

The Director, RIMS, Ranchi will appoint the Chairperson and all the committee members based on their competence, experience and integrity. Members will confirm their acceptance to the Dean by providing all the required information for membership. The Chairperson will furnish any information or report to the Dean of Faculty, RIMS, Ranchi when required.

6. COMPOSITION OF IEC RIMS, RANCHI

The EC shall be multidisciplinary and multi-sectorial in composition. The **Institution Rajendra Institute of Medical Sciences, Ranchi** shall constitute the EC. Independence and competence shall be the characteristics of EC. The minimum number of members in the committee shall be seven and maximum number will be 15. It shall be constituted keeping in mind the representation of gender, scientific and non-scientific disciplines, clinical and non-clinical disciplines, the lay community, legal expertise, social science and others to represent different points of view, and to safeguard the interests and welfare of all sections of the community / society. The Committee shall comprise of a Chairperson, a vice chairperson, a Member Secretary, a joint member secretary, and other members from the critical

categories, complying with the provisions of New Drugs and Clinical Trials Rules, 2019, and/or the National Ethical Guidelines for Biomedical and Health Research involving Human Participants of ICMR, 2017.

The composition shall be as follows:

- 1. Chairperson (not affiliated to the Institute)
- 2. Vice chairperson (not affiliated to the Institute)
- 3. Member secretary (from the Institute)
- 4. Joint Member secretary (from the Institute)
- 5. Basic Medical Scientist
- 6. Clinician
- 7. Legal expert
- 8. Lay person from the community
- 9. Social Scientists / NGO Representatives / Philosophers / Ethicist / Theologians

Expert Member/ Independent Consultants- Subject experts shall be invited to offer their views on review of research protocols and causality assessment for SAE. Their inputs shall be maintained on record and considered when reaching a decision. An expert member means a member who is a 'health care professional' (as mentioned below and registered by their respective council) and has professional qualifications or experience relating to the conduct of, or use of statistics in clinical research, unless those professional qualifications or experience relate only to the ethics of clinical research or medical treatment.

7. APPOINTMENT AND MEMBERSHIP REQUIRMENTS

- a. The Head of the Institution of Rajendra Institute of Medical Sciences, Ranchi is responsible for making the appointment of committee members.
- b. The member secretary shall be appointed from the institute.
- c. All the members will serve for a period of 3 years. The membership will be renewed after the stated term of three years.
- d. Members must accept the appointment in writing
- e. Members will be selected in their personal capacities, based on their interest, ethical and/or scientific knowledge and expertise, as well as on their commitment and willingness to volunteer the necessary time and effort for the EC's work.
- f. Members must disclose in writing any **conflict of interest**. The EC shall decide the extent to which members that might have a conflict of interest may participate in bringing out an advice/decision; refer to SOP- Confidentiality and Conflict of Interest Agreement. Members shall be required to sign a confidentiality agreement at the start of their term. [Annexure 4, 5]

7.1 EDUCATIONAL REQUIREMENTS FOR THE MEMBERS OF IEC, RIMS, RANCHI

- a. IEC members have a need for initial and continued education regarding the ethics and science of biomedical research. All IEC members must be conversant with ICMR Guidelines for Research involving Human Subjects 2006, and ICH-GCP guidelines.
- b. IEC members will receive introductory training material in research bioethics and functioning of IEC and will be exposed to ongoing opportunities for enhancing their capacity for ethical review.
- c. Training of the IEC members in Research Bioethics A new member will be inducted 1 month prior and will be requested to be an 'observer' for the first board meeting.
- d. An introductory training will be imparted by the Member Secretary.
- e. The IEC members will be encouraged to receive ongoing training by attending workshops at least once every year.
- f. The IEC will conduct workshops from time to time to impart training to the IEC members and Institutional faculty members. The training programs would be scheduled and spread over the year.

7.2 RENEWAL OF MEMBERSHIP

- a. The membership will be renewed after the stated term of three years.
- b. Selection of members shall be done at least one month in advance.
- c. Designated members of the EC who wish to attend EC meetings as observers shall read, understand, accept and sign the agreement contained in the Confidentiality / Conflict of Interest form (Annexure) at the beginning of the EC meeting and/or before scientific and ethical review tasks of the EC commence.

7.3 RESIGNATION

- a. If any member wishes to discontinue from the EC, he/she would be required to inform the Chairperson, in writing.
- b. Members may voluntarily resign from the committee at a month's notice citing appropriate reasons and incase of internal members their membership would be considered withdrawn, if they resign from the Institute.

7.4 TERMINATION/ DISQUALIFICATION PROCEDURE

During the tenure, Chairperson shall have the authority to terminate/ disqualify any of the members in the event that the member has not complied with the conditions of appointment, is absent without prior information for three consecutive meetings or on an occurrence of any event that casts a serious doubt on the integrity or ethics of the member.

In all such situations/ circumstances, the Head of Institute shall be informed of such termination to the member prior or within 15 calendar days of termination. Documentation of the termination

shall be recorded in the minutes of the next duly constituted EC meeting and the EC membership roster and circulars shall be revised.

8. OFFICE BEARERS AND MEMBER SPECIFIC ROLES AND RESPONSIBILITIES

The IEC will have the following office bearers who have the expertise and professional qualifications to review the proposals submitted.

i. Chairperson and Vice Chairperson

- The IEC, RIMS, Ranchi Chairperson should be a highly respected individual from outside RIMS, fully capable of managing the IEC and the matters brought before it, with fairness and impartiality.
- The task of making the IEC a respected part of the institutional community will fall primarily on the shoulders of this individual. The IEC must be perceived to be fair and impartial, immune from pressure either by RIMS administration, the investigators whose protocols are brought before it, or other professional and non-professional sources.
- The IEC Chairperson will respect the diverse backgrounds, perspectives, and sources of expertise of all IEC members, especially the contributions of the non-scientists, and must have the ability to foster such respect among the IEC members.
- The chairperson will ensure active participation of all members (particularly non-affiliated, non-medical/ non- technical) in all discussions and deliberations and ratify minutes of the previous meetings.
- In the absence of the chairperson of IEC for scheduled IEC meeting, Vice Chairperson will
 act as Chairperson. In case of anticipated absence of both Chairperson and Vice
 Chairperson at a planned meeting, the Chairperson should nominate a committee member
 as Acting Chairperson or the members present may elect an Acting Chairperson on the day
 of the meeting.
- The Acting Chairperson would be a non-affiliated person and will have all the powers of the Chairperson for that meeting.
- The chairperson will seek COI declaration from members and ensure quorum and fair decision making.
- He will also handle complaints against researchers, EC members, conflict of interest issues and requests for use of IEC data, etc.

ii. Member Secretary and Joint Member Secretary

- The Member Secretary will be a staff member of RIMS, committed to the task of coordinating and managing the activities of the committee.
- He/she will organize an effective and efficient procedure for receiving, preparing, circulating and maintaining each proposal for review
- He will schedule EC meetings, prepare the agenda and minutes
- The member secretary will be responsible for EC documentation, communication and archiving
- He/she will also ensure training of EC secretariat and EC members
- He/she will ensure SOPs are updated as and when required also ensure the adherence of EC functioning to the SOPs
- The member secretary will be responsible for the preparations for and audits and inspections He/she will also ensure completeness of documentation at the time of receipt and timely inclusion in agenda for EC review.
- Assess the need for expedited review/ exemption from review or full review.

iii. The Secretariat

The Secretariat will be composed of the Member Secretary, IEC, and the administrative supporting staffs. The supporting staff consists of staff members of RIMS, appointed by the Director, RIMS. The secretariat shall have the following functions:

- Organization of an effective and efficient tracking procedure for each proposal received.
- Preparation, maintenance and distribution of study files.
- Organization of regular IEC meetings.
- Preparation of the agenda and the minutes of the meetings,
- Maintenance of the IEC records and archives.
- Communication with IEC members and PIs.
- Arrangement of training for personnel and IEC members.
- Provision of the necessary administrative support for IEC related activities to the Member Secretary, IEC.
- Receipt of IEC processing fees for pharma-funded projects and the issue of official receipts for the same.

iv. The IEC Administrative Staffs

- 1. There will be administrative attendant/s /helper/s who will help the IEC Chairperson and Member Secretary in executing functions of the IEC. Additional staffs may be appointed and duties assigned as and when deemed necessary by the IEC. The eligibility criteria for new staff to be appointed will be laid down depending on the required job profile. The need for appointment of administrative staff, job profile and qualifications may be recommended by IEC members during regular IEC meeting and will be recorded in minutes. These will be forwarded to the Director, RIMS.
- 2. The administrative staff will be appointed by conducting formal interviews as per RIMS policy.

Duties of the Administrative staffs:

- Organizing an effective and efficient tracking procedure for each proposal received.
- Preparing, maintaining and distributing study files
- Organizing IEC meetings regularly
- Preparing the agenda and minutes of the meetings
- Constitution of Institutional Ethics Committee,
- Maintaining IEC records and archives.
- Communicating with IEC members and PIs.
- Arranging training for personnel and IEC members
- Providing necessary administrative support for IEC related activities to the Member Secretary, IEC.
- Receiving IEC processing fees and issuing official receipts for the same.
- Corresponding with the IEC members, external experts and investigators.
- Making the pre and post-arrangements of IEC meetings.
- Preparing the agenda and minutes of the IEC meetings.
- Answering queries of the investigators.
- Filing study related documents.
- Archiving and maintaining the study files.
- Preparation for accreditation, audits
- Training for investigators, key study personnel, IEC members, and IEC staff.
- Participate in the development and subsequent implementation of SOPs
- Developing an effective and efficient tracking procedure

Duties of the Attendant/s /Helper/s:

- a. Assisting the secretariat in arranging the IEC meetings.
- b. Dispatching sets of study documents to IEC members and external experts.

- c. Receiving the study related documents from and dispatching the IEC letters to the investigators.
- d. Filing study related documents.
- e. Archiving and maintaining the study files
- f. Corresponding with the IEC members and external experts. The IEC staff will report to the Member Secretary and/or Chairperson.
- g. The office timings for the IEC staff will be as per RIMS rules and regulations. The staff will avail leave as per RIMS norms.

v. The IEC Members

The IEC members will have following responsibilities:

- Determining the scientific and ethical validity of the research as well as the protection of the safety, rights and confidentiality of the research participants.
- Scientific and ethical review with special emphasis on the intervention, benefit-risk analysis, research design, methodology and statistics, continuing review process, SAE, protocol deviation, progress and completion report
- For clinical trials, the pharmacologist member will review the drug safety and pharmacodynamics.
- The clinician member to review the protocol (SAE, protocol deviation or violation, progress and completion report), medical care, facility and appropriateness of the principal investigator, and provision for medical care, management and compensation.
- The legal expert member will review the MoU, Clinical Trial Agreement (CTA), regulatory approval, insurance document, other site approvals, researcher's undertaking, protocol specific other permissions, such as, stem cell committee for stem cell research, HMSC for international collaboration, genomic research, compliance with guidelines etc. in the proposals and interpretation and information to EC members about new regulations if any.
- Review progress reports and monitor ongoing studies.
- Maintain confidentiality of the documents and deliberations of the IEC meetings.
- Declare conflict of interest, if any IEC members shall disclose to the IEC all conflicts of the IEC member, their spouse/domestic partner, and their dependent children. Such disclosures shall be sufficiently detailed to allow the IEC Administration to transfer the project to another IEC member or allow time for an alternate member to attend the IEC meeting to meet quorum.
- Carry out work delegated by the Chairperson, Vice-Chairperson and/or Member Secretary.
- Participate in continuing education activities in biomedical ethics and biomedical research.
- Provide information and documents related to training obtained in biomedical ethics and biomedical research to the IEC secretariat.

Directors, Head of Institution, Superintendents, Administrative officers who are responsible for business development will not be involved in the review process. In the absence of the Chairperson, Vice Chairperson will chair the meeting. In the absence of both, a member who is independent of the institution will chair the meeting as the Acting Chairperson.

9. QUORUM REQUIREMENTS

Minimum of 50% of committee strength and not less than 7 members are required to constitute the quorum for the meeting of which at least one member has to be from outside the institution, and one member will be a non-scientific member & one from opposite gender. All decisions will be taken in meetings and not by circulation of project proposals.

Quorum will have besides the Chairperson and the Member secretary, 6 members with following representations:

- a. Basic medical scientists (preferably one pharmacologist).
- b. Two clinicians
- c. Legal expert
- d. Social scientist / representative of non-governmental voluntary agency / philosopher / ethicist / theologian or a similar person
- e. Lay person from the community

10. HONORARIUM, FEES AND IEC OFFICE EXPENSES

The members of the IEC, Rajendra Institute of Medical Sciences, Ranchi shall be paid Rs 1000/-as honorarium for attending the IEC meetings and reviewing the proposals.

COMPENSATION AND REIMBURSEMENTS TO EXTERNAL MEMBERS

All external members, and experts invited (if any) will be paid an honorarium of Rs. 1000/- for each meeting attended and transport facilities would be either provided by the institution or reimbursement will be done for travel costs incurred towards contributing to the workings of the IEC according to the Institution 's norms. Appropriate bills shall have to be submitted together to the Member Secretary.

EC REVIEW AND ARCHIVAL FEE

The Ethics Committee (EC) shall charge an application fee for sponsored research projects. All research proposals/clinical trials funded/sponsored by Pharmaceutical companies, Agencies, Multinationals etc. will be charged an administrative fee/ processing fee.

Waiver of these fees is permissible for non-funded studies, departmental studies, and studies funded by organizations like ICMR, UGC, DBT, DST Government of India, State Science & Technology Department, UNICEF, WHO, USAID, Non profitable organizations etc.

All applications need to be mandatorily accompanied by application fee before it can be processed. The fee shall be paid by cheque or by demand draft drawn in favor of EC and accounts thus maintained.

INITIAL REVIEW FEE

The EC shall charge a non-refundable, initial one-time review fee as administrative charges given below:

Pharmaceutical Industry and Contract Research Organisation (CRO) Funded Clinic	al Trials
	of the budget
whichever is more.	
Investigator Initiated Projects (Funded by Non-Govt. Funding Agency)	Rs. 20,000/-
Investigator Initiated Projects (From outside RIMS)	Rs. 25,000/-
Investigator Initiated Projects (Funded by Govt. Funding Agency)	. Rs. 5,000/-
Student research (thesis)	. Rs. 500/-

STUDY RENEWAL FEE

The EC shall charge a yearly fee (Rs. 5000/) for ongoing review of the study from the second year. The study renewal review fee funds the costs of the Committee renewal review of the ongoing review of adverse events, protocol variances and site visits. The committee examines each Investigator's progress reports and activities for the previous year.

AMENDMENT FEE

The EC will charge an amendment fee of Rs 2000 for any amendment(s) in the ongoing study.

ARCHIVAL FEE

The EC will charge an amount of Rs 75,000 as archival fee for a tenure of 5 years.

All applications need to be mandatorily accompanied by the application fee before it can be processed. The fees shall be deposited by Demand Draft in favour of IEC, RIMS, payable at SBI, RMCC Branch, Ranchi or by NEFT in IEC current Account in SBI Account No. 39636660706 RMCC Branch, IFSC code no. SBIN000001672.

OFFICE EXPENSES

For the maintenance of the office, a sum of Rs 2000/- per month will be given to the secretariat.

11. APPLICATION PROCEDURE

- 1. All proposals should be submitted on any working day 1 month in advance of scheduled meeting in the prescribed Guidelines for submission of projects [Annexure No. 12]. The SOP is available on the RIMS website.
- 2. All relevant documents should be enclosed with the application form as provided in the check list [Annexure no. 14].
- 3. Required number of copies of the proposal along with the application and documents in prescribed format duly signed by the Principal Investigator (PI) and Co-investigators/ Collaborators / Research Scholars shall be guided to the Chairperson, RIMS, Ranchi, through member secretary. In his absence via any person nominated by Chairperson, receipt of the application will be acknowledged by the IEC office. The investigators submitting the projects to the IEC for the approval will submit the project according to the prescribed guidelines, summary proforma and the checklist for submission. [Annexure 12, 13 & 14] along with the subject information sheet and consent form [Annexure 9].

Every application will be allotted an IEC registration number to be used for all future correspondence and reference. The date of IEC meeting will be intimated to the PI to attend the meeting and to make a brief presentation of the proposal and to clarify the points raised by the members. The decision of the committee on the proposal will be communicated in writing. If revision is to be made, the revised document in required number of copies should be submitted within a stipulated period of time as specified in the communication or before the next meeting.

12. REVIEW PROCEDURES

- 1. The meetings of the IEC, RIMS, Ranchi will be held on periodic intervals, 2nd week of every month unless otherwise specified by the member secretary. Additional review meetings can also be held with short notice as and when required.
- 2. Additional meetings will be planned in accordance with the need for the **Expedited IEC Approval**. An expedited review (which may involve less waiting time for IRB approval) may be carried out by the IRB chairperson or by one or more experienced IRB members designated by the chairperson. The reviewers may exercise all of the authorities of the IRB except that of disapproving the research. A proposal submitted for expedited review may be disapproved only by the full IRB.
- 3. The proposals should be sent to the IEC at least 1 month in advance of scheduled meeting.
- 4. Required number of copies of the proposal along with the application and documents in prescribed format duly signed by the Principal Investigator (PI) and Co-investigators/Collaborators / Research Scholars shall be guided to the Chairperson, RIMS, Ranchi,

- through member secretary. Receipt of the application will be acknowledged by the IEC office.
- 5. The notice of each IEC meeting along with the agenda shall be sent to all the members at least one week before the meeting.
- 6. The IEC'S member-secretary shall screen the proposals for their completeness and depending on the risk involved categorize them into three types, namely, exemption from review, expedited review and full review.
- 7. Decisions will be taken by consensus after discussion, and whenever needed voting will be done. Decision of chairperson will be final.
- 8. Every application will be allotted an IEC registration number to be used for all future correspondence and reference. The date of IEC meeting will be intimated to the PI to be available and to clarify the points raised by the members if any.
- 9. The decision of the committee on the proposal will be communicated in writing. If revision is to be made, the revised document in required number of copies should be submitted within a stipulated period of time as specified in the communication or before the next meeting.
- 10. Researchers will be invited to offer clarifications if need be. The PI / research scholar will then present the proposal in person in the meeting. When the PI is not available due to unavoidable reasons the co-PI will present the proposal.
- 11. Independent consultants/experts will be invited to offer their opinion on specific research proposals if needed.
- 12. Minutes of the meeting will be written down and chairperson's approval will be taken in writing.

The IECs Member Secretary or the Secretariat shall screen all the proposals for their completeness and depending on the risk involved, categorize them into three types: Exemption from Review, Expedited Review and Full Review.

12.1 Types of Review

12.1.1 Exemption from review:

The proposals which present less than minimal risk will be exempted from the review process as may be seen in the situations like –

Research conducted on data available in the public domain

Research on educational practices

Observation of public behavior when information is recorded without any linked identifiers and disclosures would not harm the interests of the observed person

Quality control and quality assurance audits in the institute etc.

12.1.2 Expedited Review:

Proposals which present no more than minimal risk to the research participants will be subjected to expedited review. The member secretary and the chairperson of the IEC or the designated member of the committee or sub-committee of the IEC will do expedited review if the proposals involve:

Minor deviations from originally approved research during the period of approval usually of one year duration

Revised proposal previously approved through full review by the IEC or continuing review of approved proposals where there is no additional risk or activity is limited to data analysis.

Research involving clinical materials like data, records, documents or specimens that have been collected for non-research or clinical purposes.

When in emergency situations like serious outbreaks or disaster the full review is not possible, prior written permission of IEC may be taken before use of the test intervention. However, such research will be approved only for pilot study or preliminary work.

Examples of the research that may be eligible for expedited review are:

- Collection of hair or baby teeth.
- Collection of external secretions, including sweat and saliva.
- Recording of data from adults using noninvasive procedures that are routinely employed in clinical practice (not including exposure to electromagnetic radiation outside the visible range, for example, x-rays or microwaves.)
- Collection of blood samples by venipuncture.
- Voice recordings made for research purposes, such as investigations of speech defects.
- Moderate exercise by healthy volunteers.
- Study of existing data, documents, records, pathological specimens, or diagnostic specimens etc.

12.1.3 Full Review:

All the research proposals presenting more than minimal risk that are not covered under exempt or expedited review will be subjected to full committee review.

Points to Stress upon while reviewing Research Protocols

The protocol would be reviewed keeping in mind the following points:

- i. Measures to protect autonomy,
- ii. Risk/benefit determinations with respect to the vulnerability
- iii. Whether vulnerable subjects are bearing unequal burden in research.
- iv. Member of the IEC who would be reviewing such protocols should be well versed
- v. With the potential harm or risk of such population participating in the study. Special justification is required for inviting vulnerable individuals to serve as research subjects and,

- if they are selected, the means of protecting their rights and welfare must be strictly adhered to.
- vi. The extent of protection afforded should depend upon the risk of harm and the likelihood of benefit. The judgment that any individual lacks autonomy should be periodically reevaluated and will vary in different situations. The central issue for the

IEC to consider is whether the potential subject's ability to exercise free choice is limited in some way.

13. REVIEW OF THE SERIOUS ADVERSE EVENTS (SAE) REPORTS AND COMPENSATION ISSUES

13.1 Review of the SAE

- The primary responsibility of the IEC is to review and address SAE and unexpected events involving risks to research participants. In addition, the committee is authorized to offer mediation under appropriate circumstances.
- The IEC should also make sure that researchers/investigators are made aware of the policies and procedures concerning reporting and continuing review requirements.

The complete SAE / unexpected events report for detailed review shall be submitted to IEC as per the data elements for reporting SAE occurring in a Clinical trial/ Bioavailability study/ Bioequivalence study. [Annexure 8]

• Notifying the IEC does not relieve the PI from his/her responsibility to notify the sponsor and regulatory authorities.

For on-site SAE Instructions for PI:

All SAEs including Deaths should be reported within 24 hours of their occurrence to IEC, Sponsor or its representative and the Licensing authority.

After due analysis of the serious adverse event including Death, shall be forwarded by the Investigator to the Sponsor, the Chairperson of the IEC, Licensing authority and the Head of the Institution where the trial has been conducted.

The Chairperson of the Expert Committee constituted by the CDSCO (in case of death SAE) within fourteen calendar days of the occurrence of the serious adverse event of death.

SAE related activities before IEC meeting:

After SAEs is received the IEC, member secretary will verify that the reports are complete, signed

and dated by the PI/Co-PI and are checked for dates and typographical errors in the SAE event description, SAE event term etc.

Actions to be taken by IEC

The Member Secretary will review the SAE Report, and an expedited meeting will be called to review and opine on the SAE. Any queries raised shall be communicated to the PI for action. After analysis, SAE (other than death) report and opinion on financial compensation would be sent to the Licensing Authority within 30 calendar days.

In case of death SAEs, the analyzed report and opinion on financial compensation would be sent to the Chairperson of Expert Committee and Licensing Authority with 30 calendar days.

For off-site SAEs

It is the PI's responsibility to submit the offsite SAEs to IEC and one copy is acknowledged and returned back.

If any queries are raised by the IEC, Member Secretary will communicate to PI by email or letters as applicable; else the Offsite SAEs are filed in the respective project files.

Depending on the trend observed by the committee, if appropriate, specific action or combination of actions will be taken. Some of those are listed below:

- 1) Note the SAE report in the IEC records if information submitted is found to be adequate.
- 2) Direct the PI to inform participants already enrolled in the study about the SAE and request them to undertake additional visits, additional procedures, additional investigations, etc. as prescribed in the amendment.
- 3) Request further follow up information or Request additional details Recommend an amendment to the protocol, the ICD, Participant information sheet, investigator brochure and/ or any other document.
- 4) Suspend certain activities under the protocol (while going on with activities intended to protect the safety, well-being of participants who have already been enrolled);
- 5) Suspend enrolment of new research participants;
- 6) Suspend the study till amendments requested for by the IEC are accepted
- 7) Suspend the study for a fixed duration of time;
- 8) Suspend the study till additional information is obtained; or suspend the study till review is completed.
- 9) Terminate the study.

13.2 Criteria for assessment of trial relatedness of injury or death or permanent disability to be related to clinical trial or bioavailability and bioequivalence study —

Any injury or death or permanent disability of a trial subject occurring during clinical trial or bioavailability or bioequivalence study due to any of the following reasons shall be considered as clinical trial or bioavailability or bioequivalence study related injury or death or permanent disability, namely:-

- (a) adverse effect of the investigational product;
- (b) violation of the approved protocol, scientific misconduct or negligence by the sponsor or his representative or the investigator leading to serious adverse event;
- (c) failure of investigational product to provide intended therapeutic effect where, the required standard care or rescue medication, though available, was not provided to the subject as per clinical trial protocol;
- (d) not providing the required standard care, though available to the subject as per clinical trial protocol in the placebo controlled trial;
- (e) adverse effects due to concomitant medication excluding standard care, necessitated as part of the approved protocol;
- (f) adverse effect on a child in-utero because of the participation of the parent in the clinical trial.
- (g) any clinical trial procedures involved in the study leading to serious adverse event.

13.3 Recommendations of payments of compensation in case of serious adverse event (SAE)

- IEC, RIMS, Ranchi will instruct the sponsors or its representatives to provide payments of financial compensation in case of injury or death in clinical trial or bioavailability or bioequivalence study of new drug or investigational new drug-
- (1) Where any death of a trial subject occurs during a clinical trial or bioavailability or bioequivalence study, the legal heir of the trial subject shall be provided financial compensation by the sponsor or its representative, who has obtained permission to conduct the clinical trial or bioavailability or bioequivalence study, in accordance with the procedure specified in rule 42.
- (2) Where permanent disability or any other injury occurs to a trial subject during a clinical trial or bioavailability or bioequivalence study, the trial subject shall be provided financial compensation by the sponsor or its representative, who has obtained permission to conduct the clinical trial or bioavailability or bioequivalence study, in accordance with the procedure specified in rule 42.
- (3) The financial compensation referred to in sub-rule (1) or sub-rule (2) shall be in addition to any expenses incurred on medical management of the trial subject.
- (4) In the event of an injury, not being permanent in nature, the quantum of compensation shall be commensurate with the loss of wages of the subject as provided in the Seventh Schedule.

- (5) The sponsor or its representative shall give an undertaking along with the application for clinical trial permission to the Central Licencing Authority to provide compensation in the case of clinical trial related injury or death for which subjects are entitled to compensation.
- (6) Where the sponsor or its representative, who has obtained permission to conduct clinical trial or bioavailability or bioequivalence study, fails to provide financial compensation, as referred to in sub-rule (1) or sub-rule (2), the Central Licencing Authority shall, after affording an opportunity of being heard, by an order in writing, suspend or cancel the clinical trial or bioavailability or bioequivalence study or restrict the sponsor including its representative, who has obtained permission to conduct clinical trial or bioavailability or bioequivalence study, to conduct any further clinical trial or bioavailability or bioequivalence study or take any other action for such period as considered appropriate in the light of the facts and circumstances of the case.

13.4 Determination of the quantum of compensation in cases of regulatory clinical trial related injury or death

1. Formula in case of clinical trial related death:

Compensation = $(B \times F \times R) / 99.37$

Where,

B = Base amount (i.e. 8 lacs)

F = Factor depending on the age of the trial subject as per Annexure 1 (based on Workmen Compensation Act)

- R = Risk Factor depending on the seriousness and severity of the disease, presence of comorbidity and duration of disease of the trial subject at the time of enrolment in the clinical trial between a scale of 0.5 to 4 as under:
- (1) 0.5 terminally ill patient (expected survival not more than (NMT) 6 months)
- (2) 1.0 Patient with high risk (expected survival between 6 to 24months)
- (3) 2.0 Patient with moderate ris
- (4) 3.0 Patient with mild risk
- (5) 4.0 Healthy Volunteers or trial subject of no risk. However, in case of patients whose expected mortality is 90% or more within 30 days, a fixed amount of Rs. 2 lacs should be given.

2. Formula in case of clinical trial related injury (other than death):

For calculation of quantum of compensation related to injury (other than death), the compensation shall be linked to the criteria considered for calculation of compensation in cases of death of the trial subject as referred to in section of this Schedule. The quantum of compensation in case of Clinical Trial related SAE should not exceed the quantum of compensation which would have been due for payment in Case of death of the trial subject since the loss of life is the maximum injury possible.

As per the definition of SAE, the following sequelae other than death are possible in a clinical trial subject, in which the trial subject shall be entitled for compensation in case the SAE is related to clinical trial.

(i) A permanent disability: In case of SAE causing permanent disability to the trial subject, the quantum of compensation in case of 100% disability shall be 90% of the compensation which would have been due for payment to the nominee (s) in case of death of the trial subject. The quantum for less than 100% disability will be proportional to the actual percentage disability the trial subject has suffered.

Accordingly, following formula shall be applicable for determination of compensation:

Compensation = $(C \times D \times 90) / (100 \times 100)$

Where:

D = Percentage disability the trial subject has suffered.

C = Quantum of Compensation which would have been due for payment to the trial subject's nominees)

in case of death of the trial subject.

(ii) Congenital anomaly or birth defect:

The congenital anomaly or birth defect in a baby may occur due to participation of anyone or both the parent in clinical trial. Following situations may arise due to congenital anomaly or birth defect.

- (a) Still birth;
- (b) Early death due to anomaly;
- (c) No death but deformity which can be fully corrected through appropriate intervention;
- (d) Permanent disability (mental or physical). The compensation in such cases would be a lump sum amount such that if that amount is kept by way of fixed deposit or alike, it shall bring a monthly interest amount which is approximately equivalent to half of minimum wage of the unskilled worker (in Delhi). The quantum of compensation in such cases of SAE shall be half of the base amount as per formula for determining the compensation for SAE resulting into death. In case of birth defect leading to sub-clause (c) and (d) of this clause to any child, the medical management as long as required shall be

provided by the Sponsor or his representative which will be over and above the financial compensation.

(iii) Chronic life-threatening disease; and

(iv) Reversible SAE in case it is resolved.

In case of clinical trial related SAE causing life-threatening disease and reversible SAE in case it is resolved, the quantum of compensation would be linked to the number of days of hospitalisation of the trial subject. The compensation per day of hospitalization shall be equal to the wage loss. The wage loss per day shall be calculated based upon the minimum wage of the unskilled worker (in Delhi). Since, in case of hospitalisation of any patient not only the patient loses his/her wage, there will be direct or indirect losses of various kind including inconvenience, wage loss of attendant, etc. The compensation per day of hospitalisation in such case shall be double the minimum wage. Accordingly, following formula shall be applicable for determination of compensation:

Compensation = $2 \times X \times X \times X$.

Factor (F) for calculating the amount of compensation

Age	Factor
Not more than	
16	228
17	227
18	226
19	225
20	224
21	222
22	221
23	219
24	218
25	216
26	215
27	213
28	211
29	209
30	207
31	205
32	203
33	201
34	199
35	197
36	194
37,	192

and so onas given in -
New Drugs and Clinical Trial Rules_2019

Where, W = Minimum wage per day of the unskilled worker N = Number of days of hospitalization

DCGI Query on Serious Adverse Events:

In potentially contentious issues, Member Secretary, IEC will inform Chairperson and Chairperson may use his/her discretion to bring it to the full board IEC meeting. The reply will be sent to DCGI with a copy of the same to Principal Investigator.

14. REVIEW OF THE STUDY COMPLETION REPORT

Before Each Board Meeting:

The Secretariat will receive 1 copy (soft and hard) of Study Completion Report from the Principal Investigator. The study completion report is expected from the investigator within 1 month of completion of the study at the site. A brief study report containing data analysis from all the centers can be submitted by the investigator once available from the sponsor. The Secretariat shall review the report for completeness before submission for the Board meeting. If necessary, the IEC secretariat will retrieve the master file from the archiving with permission of the Member Secretary. The Secretariat shall verify the submitted Study Completion Report along with Study Completion Report to the Chairperson. Prior to sending the Study Completion Report to the Chairperson, the Secretariat will prepare the Study Completion statement and attach this also to the packet sent to the Chairperson.

The Chairperson and the Member Secretary will review the report, Study Completion Report Form and Study Completion statement and notify it to the other IEC members at the forthcoming full board meeting or the Chairperson can designate two other IEC members to review the Study report and related documents. If deemed necessary, the Chairperson may keep the report for discussion at the forthcoming IEC meeting.

The Secretariat will send the Study Completion Report Form and Study Completion statement to the designated IEC members if required. The Secretariat shall include the Study Completion Report Form in the agenda for IEC members for discussion at the full board meeting.

During the Board meeting

The Secretariat shall request the IEC member(s) designated the task to review a copy of the Final Report to present his/her comments. The Member Secretary entertains any discussion of the study. If appropriate to the discussions, the Chairperson may call for voting for final decision or whether to request further information or to take other action with the investigator.

After the Board meeting

The Secretariat will note the decision in the meeting minutes and the study shall be considered as closed if decision by IEC is "Noted". The IEC decision is notified to the investigator as a) noted in the IEC records

b) request for additional information / clarification

The Secretariat will accept and file the Final Report and get the Study Completion Report Form signed by the Chairperson and file it. With the permission of the Member Secretary, the secretariat will retrieve the file from the archiving. The final report will be placed in the master file and kept in the archival area. The Administrative Officer will archive the entire study protocol for a period of 5 years from the date of completion of the project if the decision is noted and closed.

15. DECISION-MAKING AND THE FORMAT TO ACCORD APPROVAL TO CLINICAL TRIAL PROTOCOL BY THE IEC, RIMS, RANCHI

- 1. Members will discuss the various issues before arriving at a consensus decision. When consensus is not arrived at, the decision will be made by voting procedure.
- 2. A member should withdraw from the meeting during the decision procedure concerning an application in case of a conflict of interest and it should be intimated to the chairperson prior to the review of the application and recorded in the minutes of the meeting.
- 3. Decision will be made only in meetings where quorum is complete.
- 4. Only the members can make the decisions. the expert consultants will only offer their opinions.
- 5. The PI will be intimated about the decision of the committee with IEC approval notice [Annexure 11]
- 6. Decision about the proposals will be communicated as Approved, Approved with Modifications or Rejected. Specific suggestions for modifications and reasons for modifications and reasons for rejection will be given. In cases of conditional decisions, clear suggestions for revision and the procedure for having the application revised will be specified.
- 7. Modified proposals will be reviewed by an expedited review through identified members.
- 8. Procedures for appeal by the researchers will be clearly defined.

16. INFORMED CONSENT:

- The IEC will ensure that the participants have been given sufficient, accurate information about the study.
- The Informed Consent document should contain all of the information that the participant needs to make an informed decision about taking part in the study [Please refer to annexures 15, 16. 17. 18].
- The participant must sign and date the informed consent document before taking part in any study procedures.
- The consent form should be written in non-technical language that participants would understand. Also, it should be written in language consistent with the participants' educational level, cultural views, and familiarity with research.
- The participant may withdraw consent and decline to participate in the study at any time before or after signing the consent document until their participation in the study is completed.
- The informed consent should state those aspects of the study/trial that are experimental, the risk and the benefits of the study/trial, the number of participants involved as well as the expected duration of the participant's involvement in the study/ trial.
- IEC will ensure that adequate provision is made to protect subjects' privacy and maintain the confidentiality of data.
- It should state the compensation and/or treatment available to the participant in the event of trial-related injury.
- It should state the anticipated expenses, if any, to the participant for participating in the study and the anticipated prorated payment, if any, to the participant for participating in the study.

17. SPECIAL CONSIDERATIONS / PROTECTION OF VULNERABLE POPULATION:

Special Considerations / Protection of Vulnerable Population will be taken care of by the IEC as there are specific concerns about specialized areas of research that require additional safeguards and protection. Vulnerable persons are those who are relatively (or absolutely) incapable of protecting their interests. More formally, they may have insufficient power, intelligence, education, resources, strength, or other needed attributes to protect their own interests.

Efforts should be made to ensure that individuals or communities invited for research be selected in such a way that the burdens and benefits of the research are equally distributed.

- 1. Research on genetics should not lead to racial discriminate.
- 2. Persons who are economically or socially disadvantaged should not be used to benefit others who are better off.
- 3. Rights and welfare of mentally challenged and mentally differently able persons who are incapable of giving informed consent or those with behavioral disorders must be protected. Appropriate proxy consent from the legal guardian should be taken after the person is well informed about the study, needs for participation, risks, and benefits involved, and the privacy and confidentiality procedures.

4. Adequate justification is required for the involvement of participants such as prisoners, students, subordinates, and employees, service personnel, etc. who have reduced autonomy as research participants, since the consent provided may be under duress or various other compelling reasons.

Individuals whose willingness to volunteer in a research study may be unduly influenced by the expectation, whether justified or not, of benefits associated with participation or of a retaliatory response from senior members of a hierarchy in case of refusal to participate may also be considered vulnerable. Examples are members of a group with a hierarchical structure, such as medical, pharmacy, dental, and nursing students, subordinate hospital and laboratory personnel, employees of the pharmaceutical industry, members of the armed forces, and persons kept in detention.

"Vulnerable" or "special" classes of subjects include as listed below

This list may not be exhaustive as there may be circumstances in which other groups are considered vulnerable, women for example, in an orthodox patriarchal society, or terminally ill cancer patients.

- pregnant women, human fetuses and neonates,
- prisoners,
- children.
- cognitively impaired persons
- Students and employees, sub-ordinates
- Minorities (as defined by national constitution and/or socio-economically backward, refugees, and such others.
- Economically and/or educationally disadvantaged AIDS/HIV positive subjects.
- Terminally ill Subjects
- Geriatric population

Special Requirements when Children are part of the Research

The following is required when children are enrolled in research:

- Children will not be involved in research that can be carried out equally well with adults.
- The purpose of the research is to obtain knowledge relevant to the health needs of children. For clinical evaluation of a new, drug the study in children should always be carried out after the phase III clinical trials in adults.
- For studies prior to phase III the drug has a therapeutic value in a primary disease of the children.
- The settings of the research should provide the child and the parents adequate medical and psychological support.
- Interventions intended to provide direct diagnostic, therapeutic, or preventive benefit for the individual child participants must be justified in relation to potential risks involved in the study and potential benefits to society.
- The risk presented by interventions not intended to benefit the individual child participant is low when compared to the importance of the knowledge that is to be gained.

- Interventions that are intended to provide therapeutic benefit are likely to be at least as advantageous to the individual child participant as any available alternative interventions.
- A parent or legal guardian of each child has given a proxy consent.
- The assent of the child should be obtained to the extent of the child's capabilities such as in the case of mature minors or adolescents, unless there is no medically acceptable alternative to the therapy provided or tested, and consent has been obtained from at least one parent or guardian.
- Such trials, unless an exception is justified, should be conducted in patients having a disease or condition for which the investigational product is intended. Participants in these trials should be particularly closely monitored and should be withdrawn if they appear to be unduly distressed.

Special Requirements when Adults can't give consent

A non-therapeutic clinical trial (i.e. a trial in which there is no anticipated direct clinical benefit to the participant) should be conducted in participants who personally give consent and who sign and date the written consent document.

Non-therapeutic clinical trials may be conducted in participants with consent of a legally acceptable representative (LAR) provided the following conditions are fulfilled:

- The objectives of the clinical trial cannot be met by means of a trial in participants who can give consent personally.
- The foreseeable risks to the participants are low.
- The negative impact on the participant's wellbeing is minimized and low.
- The clinical trial is not prohibited by law.
- The opinion of the CREC is expressly sought on the inclusion of such participants, and the
- written opinion covers this aspect.
- Such trials, unless an exception is justified, should be conducted in patients having a disease or
- condition for which the investigational product is intended. Participants in these trials should be particularly closely monitored and should be withdrawn if they appear to be unduly distressed.

Special Requirements when the research participants are Pregnant women or Nursing Mothers

The following is required when Pregnant or nursing women are enrolled in research:

Pregnant or nursing women should under no circumstances be the participant of any
research unless the research carries no more than minimal risk to the fetus or nursing infant
and the object of the research is to obtain new knowledge about the fetus, pregnancy and
lactation.

- As a general rule, pregnant or nursing women should not be participants of any clinical trial except such trials are designed to protect or advance the health of pregnant or nursing women or fetuses or nursing infants, and for which non-pregnant women or non-nursing mothers would not be suitable participants.
- The justification for participation of these women in clinical trials would be that they should not be deprived arbitrarily of the opportunity to benefit from investigations, drugs, vaccines or other agents that promise therapeutic or preventive benefits. Example of such trials are to test the efficacy and safety of a drug for reducing perinatal transmission of HIV infection from mother to child, trials for detecting fetal abnormalities and for conditions associated with or Aggravated by pregnancy etc.
- Women should not be encouraged to discontinue nursing for the sake of participation in research and in case she decides to do so, harm of cessation of breast-feeding to the nursing child should be properly assessed except in those studies where breast feeding is harmful to the infant. Compensation in terms of supplying supplementary food such as milk formula should be considered in such instances.
- Research related to termination of pregnancy: Pregnant women who desire to undergo Medical Termination of Pregnancy (MTP) could be made participant for such research as per The Medical Termination of Pregnancy Act, GOI, 1971.
- Research related to pre-natal diagnostic techniques: In pregnant women such research should be limited to detect the fetal abnormalities or genetic disorders as per the Prenatal Diagnostic Techniques (Regulation and Prevention of Misuse) Act, GOI, 1994 and not for sex determination of the fetus.

Special Requirements Concerning the Consent of Prisoners

- In case of prisoners as the research participants, the IEC, RIMS, Ranchi will approve the study as prisoner research.
- It will also include a prisoner advocate in its membership.
- The IEC members (exclusive of prisoner members) must have no association with the prisoner(s) involved in the research, apart from their membership with IEC.

18. WAIVER OF INFORMED CONSENT

It is the responsibility of the IEC to review and approve a request for verbal/written consent waiver. The Member Secretary will record the decision in the minutes and in the Application Form. The Chairperson will sign and date letter conveying the decision. When a request for waiver of consent is received from the Principal Investigator (PI) to the IEC, the following steps are taken:

- The IEC Secretariat will check if the concerned documents are filled completely and the required list of documents is enclosed.
- The IEC members will review the request taking into consideration the types of studies for which waiver of consent may be granted.
- The IEC will ensure that there are adequate mechanisms described in the protocol for protection of the identity of the research participants and maintaining confidentiality of the study data. (This is necessary as the participant cannot be assured directly about confidentiality of health data through a formal informed consent process, when consent waiver is granted).
- The final decision whether to grant the waiver is taken at a full board meeting unless the project is considered under expedited review

The decision regarding approval / disapproval of waiver is informed to the Principal Investigator in writing. If the waiver is not granted, the IEC will provide reasons for the same.

19. RESPONSIBILITIES OF THE SPONSORS AND THE INVESTIGATORS TO THE IEC, RIMS, RANCHI

Responsibilities of the Sponsor(s)

- i. The clinical trial Sponsor is responsible for implementing and maintaining quality assurance systems to ensure that the clinical trial is being conducted and data generated, documented and reported in compliance with the protocol and Good Clinical Practice (GCP) Guidelines issued by the Central Drugs Standard Control Organization, Directorate General of Health Services, Government of India as well as with all applicable statutory provisions. Standard operating procedures should be documented to ensure compliance with GCP and applicable regulations.
- ii. Sponsors are required to submit a status report on the clinical trial to the licensing authority at prescribed intervals.
- iii. In case of studies prematurely discontinued for any reason including lack of commercial interest in pursuing the new drug application, a summary report should be submitted within 3 months. The summary report should provide a brief description of the study, the number of patients exposed to the drug, dose and duration of exposure, details of adverse drug

- reactions, if any, and the reason for discontinuation of the study or non-pursuit of the new drug application
- iv. Any report of serious adverse event of death occurring in clinical trial, after due analysis shall be forwarded by the sponsor to chairman of the ethics committee and chairman of the expert committee constituted by the licensing authority as defined in New Drugs and Clinical Trials, 2019 with a copy of the report to the Licensing authority and the head of the Institution where the trial has been conducted within ten calendar days of occurrence of serious adverse event of death. The report of the serious adverse event other than death, after due analysis, shall be forwarded by the sponsor to the Licensing authority, Chairman of the Ethics Committee and the head of the Institution where the trial has been conducted within ten calendar days of occurrence of the serious adverse event.
- v. In case of injury or death occurring to the clinical trial subject, the sponsor (whether a pharmaceutical company or an Institution) or his representative, whosoever had obtained permission from the Licensing Authority for conduct of the clinical trial, shall make payment for medical management of the subject and also provide financial compensation for the clinical trial related injury or death in the manner as prescribed in Appendix XII of gazette notification dated 30th January 2013.
- vi. The sponsor (whether a pharmaceutical company or an institution) or his representative, whosoever had obtained permission from the Licensing Authority for conduct of the clinical trial, shall submit details of compensation provided or paid for clinical trial related injury or death, to the Licensing Authority within thirty days of the receipt of the order of the Licensing Authority.

Responsibilities of the Investigator(s):

- 1) Investigators will ensure that the IRB receives all the documents it requires to review the proposed research.
- 2) They will admit no participant to a study before the IRB has issued its written approval of the study.
- 3) The Investigator(s) shall be responsible for the conduct of the trial according to the protocol, New drugs and Clinical Trial, 2019 and the GCP Guidelines.
- 4) The investigator will report promptly to the IRB:
 - i. In case of changes to or deviations from the protocol, including changes made to eliminate immediate hazards to study participants.
 - ii. Changes that increase the risk to participants or significantly affect the conduct of the study.
 - iii. All adverse drug reactions that are both serious and unexpected.
 - iv. New information that may adversely affect the safety of participants or the conduct of the study.

- 5) Standard operating procedures are required to be documented by the investigators for the tasks performed by them.
- 6) The researchers will be accountable for ensuring the protection of the environment and resources at all the stages of the research, in compliance with existing guidelines and regulations.
- 7) During and following a subject's participation in a trial, the investigator should ensure that adequate medical care is provided to the participant for any adverse events. Investigator(s) shall report all serious and unexpected adverse events to the Licensing Authority as per the recommended guidelines, the sponsor or his representative, whosoever had obtained permission from the licensing authority for conduct of the clinical trial, and the ethics committee that accorded approval to the study protocol, within twenty-four hours of their occurrence.
- 8) The report of the serious adverse event of death, after due analysis shall be forwarded by the investigator to Chairman of the ethics committee and Chairman of the Expert Committee constituted by the Licensing authority under Appendix XII with a copy of the report to the Licensing Authority and the head of the institution where the trial has been conducted within ten calendar days of occurrence of the serious adverse event of death.

The report of the serious adverse event other than death, after due analysis shall be forwarded to the Licensing Authority, Chairman of the Ethics Committee and the Head of the Institution where the trial has been conducted within ten calendar days of occurrence of the serious adverse event.

The investigator shall provide information to the clinical trial subject through informed consent process about the essential elements of the clinical trial and the subject's right to claim compensation in case of trial related injury or death. He shall also inform the subject or His/ Her nominee(s) of their rights to contact the sponsor or his representative whosoever had obtained permission from the Licensing Authority for conduct of the clinical trial for the purpose of making claims in the case of trial related injury or death.

20. MAINTENANCE OF RECORDS FOR CLINICAL TRIALS BY THE OFFICE OF IEC, RIMS, RANCHI FOR THE CLINICAL TRIALS

- (1) The Ethics Committee shall maintain data, record, registers and other documents related to the functioning and review of clinical trial or bioavailability study or bioequivalence study, as the case may be, for a period of five years after completion of such clinical trials.
- (2) The Ethics Committee shall maintain the following records for a period of five years after completion of every clinical trial or bioavailability study or bioequivalence study, namely-
- (i) The constitution and composition of the Institutional Ethics Committee,

- (ii) The curriculum vitae of all members of the Institutional Ethics Committee, RIMS, Ranchi,
- (iii) The standard operating procedures followed by the Institutional Ethics Committee, RIMS, Ranchi,
- (iv) National and international guidelines followed by the Institutional Ethics Committee, RIMS, Ranchi,
- (v) Copies of the protocol, data collection formats, case report forms, investigators brochures, etc., submitted for review,
- (vi) All correspondence with committee members and investigators regarding application, decision and follow up,
- (vii) Agenda of all Ethics Committee meetings and minutes of all Ethics Committee meetings with signature of the Chairperson,
- (viii) Copies of decisions communicated to applicants,
- (ix) Records relating to any order issued for premature termination of study with a summary of the reasons thereof.
- (x) Recommendation given by Ethics Committee for determination of compensation,
- (xi) Records relating to the serious adverse event, medical management of trial subjects and compensation paid.
- (xii) One soft of copy of research proposals will be archived and rest of the copies will be destroyed after one year.
- (xiii) Only authorized person will have access to data related to IEC-RIMS Ranchi.

The Ethics Committee shall furnish the information maintained as and when required by the Central Licencing Authority or any other officer authorised on its behalf.

20. REFERENCES

- 1. ICMR Ethical Guidelines for Biomedical Research in Humans
- 2. CDSCO-GCP
- 3. WHO Standards and Operational Guidance for Ethics Review of Health-Related Research with Human Participants.
- 4. AIIMS-P SOP 2013-2014
- 5. TMC IEC SOP 2016
- 6. AIIMS Bhuvneshwar IEC SOP 2012-13
- 7. Office of the Human Research Protection, US Department of Health and Human Services (HHS).
- 8. CREC-STM SOP Version 3
- 9. New Drugs and Clinical Trials Rules, 2019 CDSCO

ANNEXURES	
Memo No	Dated
APPOINTMENT ORDER OF MEMI	BER OF IEC
То,	
Dr./ Mr. / Mrs.:	
Subject- Appointment as Member of Ethics Committee	
I understand that you were approached by my office for extending the state of the s	- · ·
It is my privilege to appoint you as a member in the cat Institutional Ethics Committee (IEC), (Human research) at Raj Ranchi (RIMS, Ranchi), w.e.f. the date of your joining. The appyears.	endra Institute of Medical Sciences
The Terms of Reference of your membership has been attache your acceptance of this offer and willingness to join with immer per the format in the attachment.	_
	Director RIMS, Ranchi.

AX1-V4/SOP01/V4

TERMS OF REFERENCE FOR MEMBERSHIP

- a. The members are appointed by the Director, RIMS, Ranchi.
- b. The members are appointed for a period of 3 years.
- c. The members should submit with the appointing authority, a brief profile of yours, with relevant information, as per the prescribed proforma.
- d. The members have to sign a Confidentiality Agreement at the start of your tenure. The confidentiality agreement protects the privacy and confidentiality of all parties whose information may be disclosed to the Committee in the course of its work.
- e. The members have to sign a Conflict of Interests Declaration Form at the start of your tenure. Such declaration is essential to decide your eligibility for membership. During the tenure of your membership if any new conflict of interests arises that might influence or bias your role as a member of the Committee, you should forthwith declare the same to the appointing authority, i.e Director RIMS, Ranchi
- f. The appointment becomes effective from the date the members receive the appointment letter and submit in writing their consent to join the Committee.
- g. On expiry of this 3 years tenure, the membership may be renewed for another term, provided the member agrees. Extension of membership beyond this tenure will be decided by the Director, RIMS, Ranchi.
- h. You are free to resign and withdraw your membership any time you wish, for which you need not have to give reasons. If you decide to resign you should send a written notification of resignation to the Director RIMS, Ranchi.
- i. The members should be sufficiently aware about their role and responsibilities as a member of the Committee in their capacity as clinician/ basic scientist/ legal expert/ NGO representative/ Lay person from the community.
- j. They should have exposure to and training experience in Good Clinical Practice (GCP) Guidelines and ICMR Biomedical Research Ethics Guidelines and their periodic amendments from time to time. You have to submit with the appointing authority, the copies of all such experience or participation certificates.
- k. During the membership tenure, the members should always try to avail the opportunities to attend the workshops and seminars to upgrade your knowledge and understanding in this area. They should expeditiously forward a copy of all such training certificates to the Member Secretary of the Committee. Besides, the Committee will also periodically hold awareness seminars or training events; they should attend them and update their knowledge and understanding in the area.
- I. They are required to act responsibly and attend the scheduled meetings of the Committee regularly, besides fulfilling the other responsibilities that are assigned to you by the Chairperson of the Committee.
- m. If a member fails to attend more than 3 consecutive meetings of the Committee, he/she may be relieved of the membership. Besides, they may be terminated of membership in case their conduct is found to be unbecoming of a member of the Committee.

I have carefully gone through all the terms and understood t	hem. I agree to comply with all.
Signature with date	Name

AX2-V4/SOP01/V4.0

Consent Letter Consent to be a member of IEC, RIMS, Ranchi

From:		
То		
	Director RIMS Ranchi	
Sub:		stitute Ethics Committee (Human Studies)
Ref: I	Letter No:	dated:
opinio be pub and fir	In response to your letter stated above, I Ranchi. I shall regularly participate in the regarding the ethical issues. I shall be wallished. I shall not keep any literature or strail review.	give my consent to become a member of IEC of the IEC meeting to review and give my unbiased willing for my name, profession and affiliation to udy related document with me after the discussion elated information confidential and shall not revear resonnel. I herewith enclose my CV.
Thank	ing you,	
Yours	sincerely,	
_	ure of the Member	
	Address: Telephone No: (Off) (Res)	email:

AX3-V4/SOP01/V4

Confidentiality Agreement Form for IEC Members, RIMS, Ranchi (Form IA)

I, Dr./ Mr./ Mrs	lling to join the Institutional Ethics CommitteeMember (mention membership category),
· -	ny information, ideas, data, discoveries, etc in ding clinical trials) that are disclosed/ revealed irtue of the membership;
 I shall consider all such information and applicable); I shall use such information for contemplate 	d confidential, privileged or proprietary (if
• I shall by no means disclose such information other than in situations where it is statutorily	ion verbally, visually or in writing, to anyone or legally permitted/bound.
Name	Signature with Stamp
Place	Date

AX4-V4/SOP01/V4

Conflict of Interest Declaration Form for IEC Members at RIMS, Ranchi (Form 1B)

the Ethics Committee (IEC, RIMS, Ranch	i), and am willing to join the Institutional Ethics
committee.	sclose in reference to my role as a member of this my role as a member of this committee; and I hereby
Name	Signature with Stamp
Place	Date

AX5-V4/SOP01/V4

Office order of constitution of IEC, RIMS, Ranchi

OFFICE ORDER	
I herewith establish and constitute an Ethics Committee of Rajendra Institute of Medical Sciences, Ranchi, to ensure a competent review of all ethical aspects of project proposal received and execute the same free from any bias and influence that could affect the objective.	1
The following members will constitute the Institute Ethics Committee (Human studies):	
1.	
2.	
3.	
4.	
5.	
6.	
7.	
The tenure of this membership will be for a period of 3 years from the date of appointment.	
Director, Rajendra Institute of Medical Sciences	,
Ranchi	

AX6-V4/SOP01V4

Proforma to be submitted to the Institutional Ethics Committee, RIMS, Ranchi (Human Studies) for MD/MS/DM/MCh/PhD Students (for thesis or Dissertation)/MBBS student projects

Kindly submit 03 copies of proforma and consent forms in 2 parts (in English and Hindi/other local language) to the Member Secretary, IEC, RIMS, Ranchi.

- 1. Title of the project:
- 2. Name and department/address of the investigator:
- 3. Name of Faculty (Guide/Co-guide) with designation & department: 4. Date of approval by Institute Research Cell:
- 5. Sources of funding
- 6. Objectives of the study:
- 7. Justification for the conduct of the study:
- 8. Methodology: It should provide details of number of patients, inclusion criteria, exclusion criteria, control(s), study design, dosages of drug, duration of treatment, investigations to be done.
- 9. Permission from Drug Controller General of India (DCGI) if applicable
- 10. Ethical issues involved in the study: less than minimal risk/ minimal risk/ more than minimal risk to the study subjects
- 11. Do you need exemption from obtaining Informed Consent from study subjects-if so give justifications
- 12. Whether Consent forms part 1 and 2 in English and Hindi/Other local language are enclosed? (if the consent form in local language is not applicable, appropriate explanations must be provided)
- 13. Conflict of interest for any other investigator(s) (if yes, please explain in brief)
- 14. Whether soft copy of the proforma (CD) has been attached?
- 15. We, the undersigned, have read and understood this protocol and hereby agree to conduct the study in accordance with this protocol and to comply with all requirements of the ICMR guidelines (2017)

Date and Signature of the Investigators

Date and Signature of the Guide

Signature of the Head of the Department

AX7-V4/SOP01V4

Proforma for submission of research proposals involving human participants for ethical approval from IEC, RIMS, Ranchi

- 1. Title of the research proposal
- 2. Name of the Principal Investigator with qualification and designation
- 3. Name of the Co-Investigator(s) with qualifications and designation
- 4. Name of the Institute / Hospital / Field area where research will be conducted
- 5. Protocol of the proposed research: (includes and not limited to) clear research objectives, rationale for undertaking the investigations in human participants in the light of existing knowledge, inclusion and exclusion criteria for entry of participants. Precise description of methodology of the proposed research, including sample size (with justification), type of study design (observational, experimental, pilot, randomized, blinded etc.), intended intervention, dosages of drugs, route of administration, duration of treatment and details of invasive procedures if any, Plan to withdraw or withhold standard therapies during the research. Plan for statistical analysis of the study. Ethical issues in the study and plans to address these issues.
- 6. Proposal should be submitted with all relevant enclosures like proforma, case report forms, questionnaires, follow-up cards, participant recruitment procedures and brochures, if any. Informed consent process, including patient information sheet and informed consent form in English and Hindi/Other local language(s). Investigator's brochure for trial on drugs/ devices/ vaccines/ herbal remedies and statement of relevant regulatory clearances. Source of funding and financial requirements for the project.
- 7. For any drug / device trial, all relevant pre-clinical animal data and clinical trial data from other centres within the country / other countries, if available.
- 8. Usefulness of the project/trial
- 9. Expected 'benefits' to volunteers / community. 'Benefits' to other categories if any.
- 10. Explain all anticipated 'risks' (adverse events, injury, and discomfort) of the project, efforts taken to minimize the 'risks'. Proposed compensation and reimbursement of incidental expenses and management of research related and unrelated injury/ illness during and after research period. Description of the arrangements for indemnity, if applicable in study-related injuries and description of the arrangements for insurance coverage for research participants, if applicable.
- 11. Agreement to report all Serious Adverse Events (SAE) to IEC, RIMS, Ranchi.
- 12. Other financial issues including those related to insurance.
- 13. An account of storage and maintenance of all data collected during the trial.
- 14. Research proposals approval by scientific advisory committee/Research Cell
- 15. For international collaborative study details about foreign collaborators and documents for review of Health Ministry's Screening Committee (HMSC) or appropriate Committees under other agencies/ authority like Drug Controller General of India (DCGI)

- 16. For exchange of biological material in international collaborative study a MoU/ Material Transfer Agreement between the collaborating partners.
- 17. Statement of conflicts of interest, if any.
- 18. Agreement to comply with the relevant national and applicable international guidelines, Good Clinical Practices (GCP) protocols for clinical trials.
- 19. All significant previous decisions (e.g., those leading to a negative decision or modified protocol) by other ECs or regulatory authorities for the proposed study (whether in the same location or elsewhere) and an indication of the modification(s) to the protocol made on that account. The reasons for negative decisions should be provided.
- 20. A statement on, probable ethical issues and steps taken to tackle the same like justification for washout of standard drug, or the use of placebo control.
- 21. Curriculum vitae of all the investigators with relevant publications in last five years.
- 22. Plans for publication of results / positive or negative / while maintaining the privacy and confidentiality of the study participants.
- 23. Any other information relevant to the study.

Signature of the Principal Investigator with date.

AX8-V4/SOP01V4

Annexure - 11

FORM TO BE FILLED BY PRINCIPAL INVESTIGATOR(PI) FOR SUBMISSION TO INSTITUTIONAL ETHICS COMMITTEE, RIMS RANCHI (FORM 2)

(For attachment to each copy of the proposal)

For office use	only							
Serial number	of IEC, RIN	MS, Ranchi						
				•				
To be filled by	Pl							
Title of Projec	et:							
Name, Designa Qualification	ntion, Depa	nrtment,]	Address, Teleph No., Mobile Email id		Number of projects already with investigator	Signatu date)an	re (with ad seal
Principal Inve	stigator							
Co- Investigato	ors							
1.								
2								
3								
4								
Please attach olimited to prev		urriculum vitae rs).	of a	ll Investigators	(wit	h subject spec	ific pub	lications
Sponsor inform	mation (Tie	ck appropriate b	ox)					
		a. Government						
1. Indian		Central		State		Institutional		

	b.	Private	;				I					
2. international	Go	overnm	ent		Private			UN ag	encies			
3. Industry	Natio	onal		Multi	national							
Contact Address of S	pons	or:										
Total Budget:												
Who will bear the cost	1.	Patien	ıt	2.	Projec	et		3.	Ex	empte	d	
of investigation / implants drugs / contrasts? 4. Other Agencies (Name)												
I Tune of Study:	oss ional		Case		Coho	ort		Clinic Trial		Re	eview	
	Sing cent	le	1		Multi cen	tre	,		Oth (Spec			
2.Status Review:		New						Revi	sed			
3.Clinical Trials: Drug	/Vac	cines/I	Device	/Her	bal Reme	dies:				•		
I. Does the stud	y inv	olve u	se of :									
D	rug				Devices	,			Vacc	ines		
				_								
Indian Systems of Medici of Medicine	ne / A	Alternate	Systen	1			Ang oth	-		N	A	
II. Is it approved a	nd m	arketed						•				
In India				U	K & Euro	pe				U	SA	
Other countries				Sı	pecify:			<u> </u>				
III. Does it involve If Yes, whether I									htained	Y	l'es .	No
If yes, Date of I		·	ouici i	xegui	atory autilo	iity S I	CIIIIS	51011 15 0	otamec		l'es	No
IV. Is it an Investig	gation	nal New	Drug	(IND)?							Nie
If yes, IND No.	:)	l'es	No
a) Investigator's	Broch	ure sub	mitted							7	Zes .	No
b) In vitro studies	data									Ŋ	l'es	No
c) Preclinical stud	dies d	lone								Y	Zes .	No
d) Clinical Study is:		Phase I			Phase II		Ph	nase III		Pha	ise IV	

e) Are you aware if this study /simi lar study being done elsewhere? If Yes, attach details	Yes	No
4. Brief description of the proposal - Introduction, review of literature, aim(s) & obfor study, methodology describing the potential risks & benefits, outcome measures, stawhether it is of national significance with rationale (Attach sheet with maximum 500)	atistical analy	
5.Subject selection:		
I. Number of Subjects:		
II. Duration of Study:		
111. Will subjects from both sexes be recruited?	Yes	No

IV.	Yes	No					
V.	7. Type of subject Volunteers Patients						
VI.	Vulnerable subjects (Tick the appropriate boxes) Yes No						
Pregna	ant women	Children	Elderly				
Foetus	S	Illiterate Handicapped					
Termin	Terminally ill Seriously ill Mentally/Challenged						
Econo	mically & socially backward	Any other					
VII.	Special group subjects (Tick the appropriate boxes)	Yes	No				
	Captives	Nurse/dependent					
	Students Institutionalised Armed forces						
Any other Staff							
6. Priv	vacy and confidentially						
I.	Study involves - Dire	ect Identifiers					
	Indirect Identifiers/coded						
	Completely anonym	nized / delinked					
11.	Confidential handling of data by	staff		Yes	No		
7. Use	of biological/ hazardous materia	als					
I.	Use of fetal tissue or abortus			Yes	No		
II.	II. Use of organs or body therapy Yes						
111.	Use of recombinant/gene therapy			Yes	No		
	If yes, has Department of Biotec DNA products been obtained?	hnology (DBT) approval	for	Yes	No		

IV.	Use of pre-existing / stored / left over samples	Yes	No			
V.	V. Collection for banking / future research					
VI.	VI. Use of ionizing radiation / radioisotopes					
	If yes, has Bhaba Atomic Research Centre (BARC) approval for Radioactive Isotopes been obtained?	Yes	No			
VII.	Use of infectious / bio hazardous specimens	Yes	No			
VIII.	Proper disposal of material	Yes	No			
IX.	Will any sample collected from the patients be sent abroad? , justify with details of collaborators	Yes	No			
a)	Is the proposal being submitted for clearance from Health Mini try 's Screening committee (HMSC) for International collaboration?	Yes	No			
b)	Sample will be sent abroad because (Tick appropriate box):					
	Facility not available in	n India				
	Facility in India in acc	cessible				

Facility available but not being accessed If so, reasons								
8.Consent:	Written		Oral A					
I. Consen	t Form : (Tick th	ne includ	ed elemen	its)				1
Understandable	language			Alternativ	ves to p	parti	cipation	
Statement that	study involves 1	research		Confident	iality of	f rec	cords	
Sponsor of stud	dy			Contact in	nforma	tion	l	
Purpose and pr	rocedures		Statement that consent is voluntary					
Risks & Disco	mforts		Right to withdraw					
Be ne fits				Consent for future use of biological material				
Compensation f	For participation		Bene fits if any on future commercialization eg. Genetic basis drug development					
Compensation	for study related	d injury						
*If written cons	ent is not obtaine	d, give re	easons:					
44 3371	****	_	PI/ Co-Pl				se/Counsellor	
11. Who w	ill obtain consen	it?	Research	staff		An	y other	
	vertising be do brochure, websit			cruitment of subjects? ttach a copy)			Yes	No
10. Risks & Bo	enefits:							

I.	Is the risk reaso	onable compared t	o the anticipated be	enefits to	**	
	subjects/ commu	nity/country?			Yes	No
11.	Is there physical / social / psychological/ discomfort?			Yes	No	
	If Yes, Minimal	or no risk				
	More than minir	num risk				
	High risk					
111.	Is there a benefit	a) to the	he subjects?			
			Direct		Indirect	
	b) Benefits to so	ciety		·		
11.	Data Monitorin	g				
I.	Is there a Data &	Safety Monitorin	ng Committee /Boar	rd (DSMB)?	Yes	No
11.	_	or reporting of adveporting is done to			Yes	No
Spon	isor		Ethics committee		DSMB	
111.	Is there a plan for	or interim analysis			Yes	No
IV. datab	pase?		naintenance of all	trial	Yes	No
10	If Yes, for how lo		inimation 9		Vac	NI a
12.	is there compe	nsation for parti	icipation?		Yes	No
If Ye	s, Monetary			In kind	d	
Speci	fy amount and type	e:				
13. Is	there compensati	on for injury?			Yes	No
If Yes	s, By sponsor			By Invest	igator	
By In	surance Company			By any o	other	
	o you have confli nancial / nonfinan				Yes	No
If '	Ves. snecify ·					

Conflict of interest for any other investigator(s) (if yes, please explain inbrief)	23	Yes 1 Yes 1 Yes 1 Yes 1 Yes 1		
15. Participant Information Sheet (mark √ if yes)		Attached English version		
		Attached Hindi version Certified that Hindi version is a true		
		translation of English version		
16.Participant Informed Consent form (mark √ if yes)		Attached English version		
		Attached Hindi version		
		Certified that Hindi version is a true		
		translation of, English version		
17. Whet her any work on this project has started or not?		$(\text{mark } \sqrt{\text{if yes}}, X \text{ if no})$		
		(Please enclose a separate certificate to		
		this effect).		
18. In case of clinical trials CTRI status				
	<u> </u>	<u> </u>		

Checklist for attached documents:	
Covering letter, through proper channel	
Project proposal – 05 Copies	
Curriculum Vitae of Investigators	
Brief description of proposal	
Patient information sheet	
Informed consent form	

Investigator's brochure for recruiting subjects	

Copy of advertisement / Information brochure

Copy of clinical trial protocol and / or questionnaire

HMSC/DCGI/DBT/BARC clearance if obtained

Undertaking that the study shall be done in accordance with ICMR and GCP guidelines

In case of multi-centre study, IEC clearance of other centers must be provided

Definite undertaking as to who will bear the expenditure of injury related to the project

In case an insurance cover is in tended, Insurance certificate must be provided (as per ICMR guidelines)

Permission to use copyrighted Questionnaire/Proforma

Investigator should provide undertaking what they will do the leftover sample tissue

Certificate / undertaking as mentioned in column 17

Others

AX10-V4/SOP01V4

FORMAT TO ACCORD APPROVAL TO CLINICAL TRIAL PROTOCOL BY THE IEC, RIMS, RANCHI.

[Proforma for research proposal involving human subjects to be submitted to the Institute Ethics Committee for approval]

To Dr
Dear Dr
The Institutional ethics committee or independent ethics committee (state name of the committee, as appropriate) reviewed and discussed your application to conduct the clinical trial
entitled "(date). The following documents were reviewed:
(a) Trial protocol (including protocol amendments), datedversion No.(s)(b) Patient information sheet and informed consent form (including updates, if any) in English or vernacular language.
(c) Investigator's brochure, dated, Version
no Proposed methods for patient accrual including advertisements etc. proposed to be
used for the purpose.
(d) Principal investigator's current Curriculum Vitae.(e) Insurance policy or compensation for participation and for serious adverse events occurring during the study participation.
(f) Investigator's agreement with the sponsor.
(g) Investigator's undertaking (enclosed).
The following members of the ethics committee were present at the meeting held on (date, time, place)
Name of each member with designation;
We (Approve / Approve with modifications /Reject) the trial to be conducted in its presented form.

The ethics committee to be informed about the progress of the study, any Serious Adverse Events (SAE) occurring during the study, any changes in the protocol and patient information or informed consent and to be provided with a copy of the final report.

Yours sincerely, Member Secretary, Institutional Ethics Committee, RIMS, Ranchi.

UNDERTAKING BY THE INVESTIGATOR

- 1. Full name, address and title of the Principal Investigator (or Investigators when there is no Principal Investigator).
- 2. Name and address of the medical college, hospital or other facility where the clinical trial will be conducted: Education, training & experience that qualify the Investigator for the clinical trial (Attach details including Medical Council registration number, or any other statements of qualifications)
- 3. Name and address of all clinical laboratory facilities to be used in the study.
- 4. Name and address of the Ethics Committee that is responsible for approval and continuing review of the study.
- 5. Names of the other members of the research team (Co-or sub-Investigators) who will be assisting the Investigator in the conduct of the investigations.
- 6. Protocol Title and Study number (if any) of the clinical trial to be conducted by the Investigator.
- 7. Commitments:
- (i) I have reviewed the clinical protocol and agree that it contains all the necessary information to conduct the study. I will not begin the study until all necessary ethics committee and regulatory approvals have been obtained.
- (ii) I agree to conduct the study in accordance with the current protocol. I will not implement any deviation from or changes of the protocol without agreement by the Sponsor and prior review and documented approval or favorable opinion from the ethics committee of the amendment, except where necessary to eliminate an immediate hazard to the trial subject or when the changes involved are only logistical or administrative in nature.
- (iii) I agree to personally conduct or supervise the clinical trial at my site.
- (iv) I agree to inform all trial subject, that the drugs are being used for investigational purposes and I will ensure that the requirements relating to obtaining informed consent and ethics committee review and approval specified in the New Drugs and Clinical Trials Rules, 2019 and Good Clinical Practices guidelines are met.
- (v) I agree to report to the Sponsor all adverse experiences that occur in the course of the investigation(s) in accordance with the regulatory requirements and Good Clinical Practices guidelines.
- (vi) I have read and understood the information in the Investigator's brochure, including the potential risks and side effects of the drug.
- (vii) I agree to ensure that all associates, colleagues and employees assisting in the conduct of the study are suitably qualified and experienced and they have been informed about their obligations in meeting their commitments in the trial.
- (viii) I agree to maintain adequate and accurate records and to make those records available for audit or inspection by the Sponsor, ethics committee, Central Licencing Authority or their authorised representatives, in accordance with regulatory provisions and the Good Clinical Practices guidelines. I will fully cooperate with any study related audit conducted by regulatory officials or authorised representatives of the Sponsor.
- (ix) I agree to promptly report to the ethics committee all changes in the clinical trial activities and all unanticipated problems involving risks to human subjects or others.

- (x) I agree to inform all serious adverse events to the Central Licencing Authority, sponsor as well as the ethics committee within twenty-four hours of their occurrence. In case, of failure to do so, I shall furnish the reason for the delay to the satisfaction of the Central Licencing Authority along with the report of the serious adverse event.
- (xi) The report of the serious adverse event, after due analysis, shall also be forwarded by me to the Central Licencing Authority, the Chairperson of the ethics committee and the Head of the institution where the trial has been conducted within fourteen days in accordance with the regulatory requirements.
- (xii) I will maintain confidentiality of the identification of all participating subjects and assure security and confidentiality of study data.
- (xiii) I agree to comply with all other requirements, guidelines and statutory obligations as applicable to clinical Investigators participating in clinical trials.
- 8. Signature of the Investigator with date.

RIMS IEC Office requires review of an approved study not less than once per six (06) months period. Therefore, a continuing review application must be submitted to the IEC in order to continue the study beyond the approved period. Failure to submit a continuing review application in a timely fashion will result in termination of the study, at which point new participants may not be enrolled and currently enrolled participants must be taken off the study.

Sincerely,

Member Secretary, IEC

AX11-V4/SOP01V4

Data Elements for Reporting Serious Adverse Events (SAE) occurring in a Clinical Trial or bioavailability or bioequivalence study

1. Patient Details

Initials & other relevant identifier (hospital/OPD record number)*

Age/DOB

Weight

Height

2. Suspected Drug(s)

Generic name of the drug*

Indication(s) for which suspect drug was prescribed or tested Dosage form and strength

Daily dose and regimen (specify units - e.g., mg, ml, mg/kg)

Route of administration

Starting date and time of day

Stopping date and time, or duration of treatment

3. Other Treatment(s)

Provide the same information for concomitant drugs (including nonprescription/OTC drugs) and non-drug therapies, as for the suspected drug(s).

4. Details of Suspected Adverse Drug Reaction(s)

Full description of reaction(s) including body site and severity, as well as the criterion (or criteria) for regarding the report as serious. In addition to a description of the reported signs and symptoms, whenever possible, describe a specific diagnosis for the reaction*

Start date (and time) of onset of reaction Stop date (and time) or duration of reaction

Dechallenge and rechallenge information

Setting (e.g., hospital, out-patient clinic, home, nursing home)

5. Outcome

Information on recovery and any sequelae; results of specific tests and/or treatment that may have been conducted

For a fatal outcome, cause of death and a comment on its possible relationship to the suspected reaction; Any post-mortem findings.

Other information: anything relevant to facilitate assessment of the case, such as medical history including allergy, drug or alcohol abuse; family history; findings from special investigations etc.

6. Details about the Investigator*

Name

Address

Telephone number Profession (specialty)

Date of reporting the event to Licensing Authority:

Date of reporting the event to Ethics Committee overseeing the site:

Signature of the Investigator

*Note: Information marked * must be provided."*

AX12/V4/SOP01V4

Institute Ethics Committee, RIMS, Ranchi Six Monthly Progress of Project

Institute Ethics Committee Reference No.	
Study title:	
Name of the Principal Investigator	
Designation / Department	
Duration of Study Date of Starting of the Study	
Progress:	
Side Effect if any:	
Amendments if any:	
Discontinuation reasons:	
Progress:	
Period of six-monthly progress report: fromto	
Signature of Principal Investigator	

AX13/V4/SOP01V4

Proforma of the Subject Information Sheet Institutional Ethics Committee (IEC) RIMS, Ranchi

Title of the project:

Site of the investigation:

Name and address of the Principal Investigator:

Contact number of Principal Investigator:

- 1. Aims and methods of the research (A brief introduction about the investigation along with purpose of the study and procedure of investigation involving human subjects in simplified manner (10-15 lines).
- 2. Expected duration of the subject participation.

 The benefits to be expected from the research to the Participants or to others.
- 3. Alternative treatment/procedure options.
- 4. Right to prevent use of biological samples (DNA, cell line etc.) at any time during the research.
- 5. Any risk to the subject associated with the study.
- 6. Maintenance of confidentiality of records.
- 7. Provision of free treatment for research related injury.
- 8. Compensation of subjects for disability or death resulting from such injury.
- 9. Freedom of individual to participate or to withdraw from research at any time without penalty or loss of benefits to which the subject would otherwise be entitled.
- 10. Amount of clinical sample in quantity, to be taken should be mentioned.
- 11. Source of funding for the Investigation.
- 12. In case of drug trials:
 - a) The chemical name of drug, date of its manufacturing and batch number must be mentioned.
 - b) Initial bio equivalent study of the drug/references should be provided
- 13. Foreseeable extent for information on possible current and future usage of the biological material and of the data to be generated from the research and if the material is likely to be used for secondary purposes or would be shared with others, clear mention of the same.
- 14. Risk of discovery of biologically sensitive information.
- 15. Publication, if any, including photographs and pedigree charts.
- 16. Responsibility of Investigators.

AX14/V4/SOP01V4

Consent Form for participants more than 18 years of age (Form 3A)

Participant Informed Consent Form

Study Title:
Study Number:
Subject's Initials:
Subject's Name:
Date of Birth/Age:

- 1. I agree voluntarily to take part in this study.
- 2. I have been explained the purpose and other details of the study. I have been given a full explanation of the procedures involved.
- 3. I have been given an opportunity to ask questions and all my questions have been answered to my satisfaction.
- 4. I am free to withdraw from the study at any time without any reason and without my medical care or legal right being affected.
- 5. I understand that the information in my medical records is essential to evaluate the results of the study. I agree to release this information on the understanding that will be treated confidentially.
- 6. I agree that I will not be referred to by name in any reports/documents/any other means concerning this study.
- 7. I have been explained the risks and benefits for the patients and society associated with the study.
- 8. I understand that if I am harmed as result of taking part in the study, treatment will be provided free of cost by the PI/Institution/University.
- 9. I agree/ do not agree that the biological samples collected during this study may be stored for future use.

I willingly agree to take part in the above study.

Note: Make 2 copies of the Subject Information Sheet and Consent Form, one copy for the Principal Investigator and the other copy for the patient.

AX15/V4/SOP01V4

Signature page for research involving children ages from birth to 6 years of age or unable to provide assent for other reasons

Parents/Legally accepted representative (LAR) Consent Form (Form 3B)

- 1. I agree that my child is voluntarily taking part in this study.
- 2. I have been explained the purpose and other details of the study. I have been given a full explanation of the procedures involved.
- 3. I have been given an opportunity to ask questions and all my questions have been answered to my satisfaction.
- 4. I know that my child is free to withdraw from the study at any time without any reason and without my medical care or legal right being affected.
- 5. I understand that the information in my child's medical records is essential to evaluate the results of the study. I agree to release this information on the understanding that will be treated confidentially.
- 6. I agree that my child will not be referred to by name in any reports/documents/any other means concerning this study.
- 7. I have been explained the risks and benefits for the patients and society associated with the study.
- 8. I agree that if my child is harmed as result of taking part in the study, treatment will be provided free of cost by the PI/Institution/University.
- 9. I agree that the biological samples collected during this study may be stored for future use

I willingly agree that my child will take part in the above study.

Allow	Do not allow
Signature of the parent/guardian Name: Age: Address:	Date:
Signature of the doctor/Principal Investigator:	Date:
Signature of the witness:	Date:

Principal Investigator and the other copy for the patient.

AX16-V4/SOP01V4

Waiver of assent
The assent of (name of child/minor) was waived because of: Age:
Maturity:
Psychological state of the child:
Signature of the Parent/Legally authorized representative: Date:
Note: Make 2 copies of the Subject Information Sheet and Consent Form, one copy for the

AX16-V4/SOP01V4

Signature page for research involving children aged 7 through 17 years of age and able to provide assent (Form 2C)

- 1. I agree that my child is voluntarily taking part in this study.
- 2. I have been explained the purpose and other details of the study. I have been given a full explanation of the procedures involved.
- 3. I have been given an opportunity to ask questions and all my questions have been answered to my satisfaction.
- 4. I know that my child is free to withdraw from the study at any time without any reason and without my medical care or legal right being affected.
- 5. I understand that the information in my child's medical records is essential to evaluate the results of the study. I agree to release this information on the understanding that will be treated confidentially.
- 6. I agree that my child will not be referred to by name in any reports/documents/any other means concerning this study.
- 7. I have been explained the risks and benefits for the patients and society associated with the study.
- 8. I agree that if my child is harmed as result of taking part in the study, treatment will be provided free of cost by the PI/Institution/University.
- 9. I agree/ do not agree that the biological samples collected during this study may be stored for future use.

I willingly agree that my child will take part in the above study.

the other copy for the patient.

Signature of the parent/guardian	Date:
Assent of child (name of child study	/minor) has agreed to participate in above
Signature of the child	Date:
Name: Age: Address:	
Signature of the doctor/Principal Investigator:	Date:
Signature of the witness:	Date:

SOP V4.0 69

Note: Make 2 copies of the Subject Information Sheet and Consent Form, one copy for the Principal Investigator and

AX17-V4/SOP01V4

Informed Consent Document for Drug Clinical Trial (FORM 2D, FORM 2E)

(As per table 03 of New Drugs and Clinical Trial Rule 2019, dated 19th March, 2019)

INFORMED CONSENT

1. Checklist of Review of informed consent documents for clinical trial subject

- 1.1 Essential elements:
- (i) Statement that the study involves research and explanation of the purpose of the research.
- (ii) Expected duration of the participation of subject.
- (iii) Description of the procedures to be followed, including all invasive procedures.
- (iv) Description of any reasonably foreseeable risks or discomforts to the Subject.
- (v) Description of any benefits to the Subject or others reasonably expected from research. If no benefit is expected Subject should be made aware of this.
- (vi) Disclosure of specific appropriate alternative procedures or therapies available to the Subject.
- (vii) Statement describing the extent to which confidentiality of records identifying the Subject will be maintained and who will have access to Subject's medical records.
- (viii) Trial treatment schedule and the probability for random assignment to each treatment (for randomized trials).
- (ix) Statement describing the financial compensation and the medical management as under:
- (a) In case of an injury occurring to the subject during the clinical trial, free medical management shall be given when required or till such time it is established that the injury is not related to the clinical trial, whichever is earlier.
- (b) In the event of a trial related injury or death, the sponsor or his representative or the investigator or centre, as the case may be, in accordance with the rule 39, as the case may be, shall provide financial compensation for the injury or death.

- (x) An explanation about whom to contact for trial related queries, rights of Subjects and in the event of any injury.
- (xi) The anticipated prorated payment, if any, to the subject for participating in the trial.
- (xii) Responsibilities of subject on participation in the trial.
- (xiii) Statement that participation is voluntary, that the subject can withdraw from the study at any time and that refusal to participate will not involve any penalty or loss of benefits to which the subject is otherwise entitled.
- (xiv) Statement that there is a possibility of failure of investigational product to provide intended therapeutic effect.
- (xv) Statement that in the case of placebo-controlled trial, the placebo administered to the subjects shall not have any therapeutic effect.
- (xvi) Any other pertinent information.
- 1.2 Additional elements, which may be required:
- (a) Statement of foreseeable circumstances under which the participation of the subject may be terminated by the Investigator without his or her consent.
- (b) Additional costs to the subject that may result from participation in the study.
- (c) The consequences of a Subject's decision to withdraw from the research and procedures for orderly termination of participation by Subject.
- (d) Statement that the Subject or Subject's representative will be notified in a timely manner if significant new findings develop during the course of the research which may affect the Subject's willingness to continue participation will be provided.
- (e). A statement that the particular treatment or procedure may involve risks to the Subject (or to the embryo or foetus, if the Subject is or may become pregnant), which are currently unforeseeable.
- (f) Approximate number of Subjects enrolled in the study.

2. Format of informed consent form for Subjects participating in a clinical trial:

Participant Informed Consent Form to participate in a Clinical Trial (Form 2E)

Study Title:		
Study Number:		
Subject's Initials:		
Subject's Name:		
Date of Birth/Age:		
Address of the Subject		
Qualification		
Occupation: Student or Self-Employed or Service or Housewife or Others (Please click appropriate) .	as	
Annual Income of the subject:		
Name and address of the nominees and his relation to the subject (for the purpose of compensation in case of trial related death).		
(i) I confirm that I have read and understood the information sheet	[-
dated for the above study and have had the opportunity to ask questions.		
(ii) I understand that my participation in the study is voluntary and that I am free to withdraw at any time, without giving any reason, without my medical calegal rights being affected.	[are or]
(iii) I understand that the Sponsor of the clinical trial, others working on the Sponsor's the Ethics Committee and the regulatory authorities will not need my permission to look health records both in respect of the current study and any further research that may be conducted in relation to it, even if I withdraw from the trial. I agree to this access.		

However, I understand that my identity will neparties or published.	ot be re	vealed in any inform	nation release	ed to third
(iv) I agree not to restrict the use of any data use is only for scientific purposes.			is study prov	ided such a
(v) I agree to take part in the above study.	[1		
Signature (or Thumb impression) of the Subject	ct/Lega	lly Acceptable Repr	resentative:	
Signatory's Name:				
Signature of the Investigator:	_		Date:	_/_
Study Investigator's Name:				
Signature of the Witness		Date:	_//	
Name of the Witness:				

Copy of the Patient Information Sheet and duly filled Informed Consent Form shall be handed over to the subject his or her attendant.

AX18-V4/SOP01V4

CHECKLIST FOR VERIFICATION OF PROPOSALS SUBMITTED TO IEC, RIMS, RANCHI [HUMAN STUDIES] (FORM4)

For official use only

Proposal No.

		Yes	No	NA	Comments
Is	all the documentation provided?				
Sci	entific importance and validity				
l.	Will the study lead to improvements in human healthand wellbeing or increase knowledge?				
2.	If the study is a replication of a previous study, is it justified?				
3.	Can the intervention studied be practically implemented?				
4.	Is there provision for dissemination of results of the research?				
5.	Has the research protocol been approved by a competent body?				
6.	Should the study be referred to a technical expert, policy marker or statistical expert? (If Yes, please inform the Secretary as soon as possible, suggesting a suitable person)				
7.	Are the objectives stated clearly?				
8.	Is the study design appropriate in relation to the objectives?				
9.	Are the investigators' qualification, competence, and experience appropriate to conduct the study?				
10.	Are the facilities at the site adequate to support the study?				
11.	Is the manner in which the results of research will be reported and published ethical?				
As	ssessment of Risk / Benefits				
I.	Is the involvement of human participants necessary to obtain the necessary information?				
2.	Are the researcher's qualifications, competence and experience suitable to ensure safe conduct of the study?				
3.	Is the justification of predictable risks and inconveniences weighed against the anticipated benefits for the research participant and the concerned committee adequately?				

		1		1	
		Yes	No	NA	Comments
	Are there any plans to withdraw or withhold standard therapy or the purpose of research and such actions if any justified?		1112		
	s there provision for compensation for participants who sustain injuries?				
6. I	Have adequate provisions been made for dealing with and reporting adverse effects?				
	Have adequate provisions been made for safety monitoring and termination of the research project?				
Resp	pect for the dignity of the research participants				
Infor	med consent				
l. I	s the process for obtaining informed consent appropriate?				
2. /	Are the pa1ticipants competent to give consent?				
	s the justification adequate for the intention to include ndividuals who cannot consent?				
4. \	Will dissent be respected?				
ŗ	s the written and oral information to be given to the research participants appropriate, adequate, complete				
	understandable? Do you approve the incentives offered?				
	s the consent given voluntarily and not due to deception, ntimidation or inducement?				
Conf	identiality	1	1	1	
	Will the researcher collect only the minimum				
	mation/ samples required to fulfil the study objectives?				
2. I	s the privacy of the research participant safeguarded?				
3. /	Are data/sample storage and disposal procedures adequate?				
Right	ts of the participants				
	s the participant's right to unconditionally withdraw from the esearch at any time safeguarded?				
2. I	s there provision for participants to be informed				
abou	t newly discovered risks or benefits during the study?				
	Is there provision for the subjects to be informed of results of clinical research?				

		Yes	No	NA	Comments
Fai	ir participant selection	1	1	•	
l.	Has the study population been determined, primarily, based on the scientific goals of the study (and not on convenience, ethnicity, age, gender, literacy, culture or economic status?				
Is the selection of participants (inclusion and exclusion criteria) appropriate so that risks are minimized and benefits are maximized and the burden of research equitably distributed?					
Does the selection of participants stigmatize any group?					
Does selection of subjects favor any group?					
5.	Is the research conducted on vulnerable individuals or groups?				
6.	Is the research externally sponsored?				
7.	Is the research a community research?				
3.	Is the research a clinical trial?				
Res	sponsibilities of the researcher				
	Is the medical care to be provided to the research participants during and after the research adequate?				
2.	Has the researcher obtained permission from the relevant authorities?				
3.	Are there any conflicts of interest, including payment and other rewards?				
4.	Are there any other/ legal/ social/ financial issues in the study?				
itio	nal Comments:	I	l	L	1

Recommendation: Approve [] Reject [] Conditional Approval (please state the conditions)	

Name of the Reviewer Signature Date

AX19-V4/SOP01V4