



### STANDARD OPERATING PROCEDURES

# POSTGRADUATE INSTITUTE OF MEDICINE UNIVERSITY OF COLOMBO

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#### **ABBREVIATIONS**

BOM - Board of Management

CIOMS - Council for International Organizations of Medical Sciences

DoH - Declaration of Helsinki

ERC - Ethics Review Committee

ICF - Informed Consent Form

ICHGCP - International Conference on Harmonization, Guidance on Good

**Clinical Practice** 

IRB - Institutional Review Board

IS - Information Sheet

MOU - Memorandum of Understanding

PGIM - Postgraduate Institute of Medicine

PI - Principal Investigator

SOP - Standard Operating Procedures

TOR - Terms of Reference

UOC - University of Colombo

WHO - World Health Organization

WMA - World Medical Association



#### SOP - 001 ERC Functions Version 2 23/04/2018



#### 1. Purpose: To describe the overall functions and scope of ERC

The Ethics Review Committee (ERC), Postgraduate Institute of Medicine, University of Colombo (PGIM/UOC) is established to safeguard the mental, physical and social well-being, dignity, rights and safety of all human subjects participating in biomedical research; and to promote standards of human research through ethical, efficient and effective review and monitoring processes in accordance with the guidelines of the Forum of Ethics Review Committees in Sri Lanka (FERCSL Guidelines) and other relevant national and international legislations and guidelines.

#### 2. Detailed instructions:

#### The Scope of Responsibilities:

- 2.1. ERC, PGIM/UOC shall
  - a. provide oversight on all matters relating to ethics of research projects involving human subjects.
  - b. ensure that the fundamental principles of research viz. autonomy, beneficence, non-maleficence and justice are adhered to in research involving human subjects.
  - c. provide independent, competent, timely ethics review and monitoring of research projects involving human subjects.
- 2.2. The ERC shall review only research proposals submitted by trainees, trainers, extended faculty and staff of Postgraduate Institute of Medicine, University of Colombo except as provided hereunder:
  - a. The ERC, PGIM/UOC may accept as valid, an ethics approval given by the ERC/IRB of another institution, for the purpose of approving the commencement of a project.
  - b. The ERC may review research proposals from researchers outside the PGIM/UOC provided a valid and current Memorandum of Understanding

(MOU) between the PGIM/UOC and the institution to which the researcher is accredited exists. Such MOU shall define:

- i. the role of the ERC in providing ethics approval and monitoring of the research;
- ii. the role of the institution to which the researcher is accredited in giving approval for the research to be conducted within its premises;
- iii. a statement indemnifying the PGIM/UOC from responsibility for liabilities that may arise from the ethics review conducted by the ERC; and
- iv. a statement that the institution to which the researcher is accredited bears responsibility for liabilities arising from the conduct of research.
- 2.3. All applications will be subject to a handling fee as decided by the Board of Management, PGIM/UOC on the recommendation of the Finance Committee, PGIM/UOC (Table 01).
- 2.4. The ERC will review research protocols in accordance with the guidelines of the Forum of Ethics review committees in Sri Lanka (FERCSL), relevant national and international guidelines and national and international legal requirements in order to determine their acceptability. This shall include an examination of the scientific and technical aspects of the proposal.
- 2.5. ERC, PGIM/UOC shall seek advice from another ERC and/or an external reviewer if the committee lacks the expertise among its members to review a specific subject or technical areas.
- 2.6. The terms 'human research projects' include, but are not limited to, research involving pharmaceuticals, medical devices, medical radiation and imaging, surgical procedures, biological samples, medical records and clinical databases, as well as epidemiological, social and psychological investigations using human subjects.
- 2.7. The ERC may review projects involving quality assurance including audits and student feedback.
- 2.8. ERC, PGIM/UOC shall not function as a committee funding research and approving research grants.
- 2.9. ERC, PGIM/UOC shall not advise clinicians regarding ethical issues which may arise in their routine clinical/medical practice.





#### SOP – 002 Membership Composition Version 2 23/04/2018

#### 1. Purpose: To describe the membership composition of the ERC

The Ethics Review Committee, Postgraduate Institute of Medicine, University of Colombo (ERC, PGIM/UOC) includes scientists and non-scientists as well as institutional and non-institutional personnel. It is independent in its reflection, advice and decision. This SOP describes the Terms of Reference (TOR) which provide the framework for the constitution of ERC, PGIM/UOC.

- 2.1. The composition of the ERC shall be in accordance with the FERCSL and other relevant national and international guidelines.
- 2.2. Membership comprises of at least eleven (11) and not more than nineteen (19) members.
- 2.3. Members shall be appointed to ensure the ERC has the expertise required to assess the applications submitted to it for consideration.
  - 2.3.1. Membership shall include the following categories:
    - a. members from PGIM
    - b. members of scientific/medical/dental institutions other than PGIM
    - c. members of scientific/medical/dental professions
    - d. members not representing scientific/medical/dental professions
    - e. a lawyer
- 2.4. Members shall be appointed to the ERC to ensure diversity including in gender, language, and culture.
- 2.5. The administrative staff of the PGIM/UOC shall not be members.
- 2.6. Where required, the ERC may seek advice and assistance from appropriate external experts to assist with the review of a protocol as detailed in SOP 011.



#### **SOP - 003 Appointment of ERC members** Version 2 23/04/2018



### 1. Purpose: To describe the procedure for the appointment of members to ERC, PGIM/UOC

This SOP describes the procedure for the appointment of members to the ERC. Members are appointed by the Director, PGIM/UOC on the recommendation of Board of Management of the PGIM as individuals for their knowledge, qualities and experience and not as representatives.

- 2.1. Members of ERC, PGIM/UOC are appointed as individuals for their knowledge, qualities and experience and not as representatives of any organisation, group or opinion.
- 2.2. Prospective members of ERC, PGIM/UOC shall be appointed by nominations or advertisement.
- 2.3. Recruitment by nomination: When the expertise of a specific individual is required the ERC will request the Director, PGIM/UOC to take steps to appoint the said individual to the ERC.
- 2.4. Recruitment by advertisement: Applications for membership shall be called from members of the Boards of Study, PGIM/UOC by the Director, PGIM/UOC.
- 2.5. Prospective members are invited to sign a confidentiality undertaking (as per Annex 1) and attend a meeting of the ERC as observers.
- 2.6. Prospective members shall provide a copy of their curriculum vitae (as per Annex 2) to the Board of Management, PGIM/UOC through Director, PGIM/UOC.
- 2.7. A selection committee, consisting of the Chairperson, the Secretary and a member of the ERC shall interview the prospective applicants and make recommendations to the Director, PGIM/UOC.
- 2.8. Letters of appointment (as per Annex 3) will be issued by the Director, PGIM/UOC. The letter of appointment shall include the date of appointment,

- length of tenure, conditions of appointment, assurance that indemnity will be provided in respect of liabilities that may arise in the course of bona fide conduct of duties as a member of the ERC, PGIM/UOC and the circumstances whereby membership may be terminated according to the relevant SOPs.
- 2.9. Members of ERC, PGIM/UOC shall agree to their names and professions being made available to the public and being published on the PGIM/UOC website.
- 2.10. Upon appointment, members shall be provided with the following documents:
  - 2.10.1. Letter of appointment (Annex 3)
  - 2.10.2. Copy of the confidentiality agreement (Annex 1)
  - 2.10.3. Standard Operating Procedures
  - 2.10.4. An updated list of ERC members including their names and contact information
  - 2.10.5. Any other relevant information about the ERC's processes, procedures, and proposals
- 2.11. Duration of membership will be for a period of three (03) calendar years.
- 2.12. Members are eligible for re-appointment. The committee shall be reconstituted at the end of three (03) years. The new committee should comprise of at least seven (07) members who have a minimum of two years' experience as members of the preceding ERC, PGIM/UOC in order to maintain the expertise and to facilitate the efficient functioning and continuity of the ERC.
- 2.13. New members are expected to attend training sessions.
- 2.14. Non-affiliated members will receive an allowance and travel expenses in accordance with the PGIM regulations.
- 2.15. Members may seek leave of absence from the ERC for extended periods and steps shall be taken to fill the temporary vacancy if the period exceeds three months.
- 2.16. Membership shall lapse if a member fails to attend three consecutive meetings of the ERC without a reasonable excuse/apology, unless exceptional circumstances exist. Such circumstances should be notified to the ERC in writing.
  - a. A valid excuse is defined as being involved in designated academic or clinical work. This should be informed to the ERC in writing prior to the

- commencement of the ERC meeting for which the member is going to be absent.
- b. The Director, PGIM/UOC will notify the members of such lapse of membership in writing. Steps shall be taken to fill the vacancy.
- 2.17. The Director, PGIM/UOC may take steps to dissolve the ERC and appoint a new committee if the ERC fails to carry out its functions to the satisfaction of the Board of Management, PGIM/UOC. The Director may take this action only if requested to do so by the majority of the members of the Board of Management, PGIM/UOC.
- 2.18. A member may resign from the ERC at any time upon giving notice in writing to the Chairperson/ERC and the Director, PGIM/UOC. The effective date of resignation will be the date on which the resignation is formally accepted by the Director, PGIM/UOC and BOM, PGIM/UOC.
- 2.19. Vacancies in the ERC will be filled as per SOP 002 and SOP 003.
- 2.20. The ERC shall elect the Chairperson and Secretary from amongst its members and inform Director, PGIM/UOC who will issue a formal notice of appointment. (An individual should have relevant training and at least three (03) years' experience in ethics review to be eligible for the posts of Chairperson and Secretary).
- 2.21. Subcommittees of Ethics Review Committee: The term subcommittee refers to a group of at least three ERC members appointed by the Chairperson, to manage a specific task or make a particular decision or recommendation relating to the functioning or standard operating procedures of ERC, PGIM/UOC.
- 2.22. Members are expected to participate in relevant specialized subcommittees as and when required.



#### SOP – 004 Responsibilities of ERC Members



Version 2 23/04/2018

#### 1. Purpose: To describe the functions of members of the ERC.

This SOP describes the responsibilities of the members of ERC, PGIM/UOC.

#### 2. Detailed instructions:

#### 2.1. Responsibilities of ERC members:

- 2.1.1. Attend meetings regularly and remain until meetings are adjourned. Those who are unable to participate in meetings in person can join the meetings through video conferencing or teleconferencing with prior notice.
- 2.1.2. Remain independent, impartial, and objective
- 2.1.3. Maintain confidentiality with regard to all matters pertaining to the ERC.
- 2.1.4. Disclose conflicts of interests and where a conflict exists, refrain from reviewing, and leave the room during deliberations and voting.
- 2.1.5. When assigned as primary reviewers;
  - a) Complete and handover assessment forms to the Secretary two (02) working days prior to the scheduled ERC meeting. If unable to attend, the forms should be sent to the Secretary ERC two (02) working days before the scheduled ERC meeting.
  - b) Lead the discussion and summarize in order to make decisions at full board meetings.
- 2.1.6. Decide by vote or consensus, whether to approve, request revisions, not approve or defer studies following deliberation at full board meetings.
- 2.1.7. Keep up-to-date with national and international research ethics and regulatory guidance.
- 2.1.8. Perform any other duties assigned to members according to the SOPs.

2.1.9. Perform any other duties assigned by the Chairperson/Secretary.

#### 2.2. Responsibilities of Chairperson:

- 2.2.1. Conduct all meetings of the ERC according to the SOPs. Provide guidance to ERC members and staff.
- 2.2.2. Perform duties assigned to the Chairperson according to the SOPs
- 2.2.3. Periodically review existing and formulate new ERC policies and guidelines in consultation with the members of ERC.

#### 2.3. Responsibilities of Secretary:

- 2.3.1. Organize meetings, maintain records and communicate with all concerned parties.
- 2.3.2. Supervise, guide and monitor office staff.
- 2.3.3. Prepare minutes of the meetings and general correspondence in concurrence with the Chairperson.
- 2.3.4. Prepare the agenda for the ERC meeting.
- 2.3.5. Ensure that membership files are current and up to date.
- 2.3.6. Assign primary reviewers for applications in consultation with the Chairperson and coordinate the review process.
- 2.3.7. Perform duties assigned to the Secretary according to the SOPs.
- 2.3.8. Perform any other duties assigned by the Chairperson.

#### 2.4. Responsibilities of Support Staff

- 2.4.1. Coordinate and process all initial, continuing review and study modification submissions.
- 2.4.2. Prepare letters to applicants, relaying specific ERC requests and follow-up.
- 2.4.3. Assist the Secretary in preparation of official minutes of the meetings.
- 2.4.4. Coordinate electronic (or other) distribution of applications and related documents received for review.
- 2.4.5. Maintain communications with members of the ERC.

- 2.4.6. Circulate relevant information, records etc. to the members of the ERC.
- 2.4.7. Maintain the electronic database of the ERC.
- 2.4.8. Perform any other duties pertaining to the ERC assigned by the Chairperson and the Secretary.



#### SOP - 005 Orientation of New Members and Training Version 2 23/04/2018



### 1. Purpose: To describe the procedure for the orientation of new members and training.

Upon being appointed to the ERC, PGIM/UOC as a new member, such members are expected to follow the instructions given in this SOP.

- 2.1. New members shall receive an electronic copy of SOPs and TORs as per SOP 003
  2.8. It is the responsibility of the members to read and understand their functions as members of the ERC, PGIM/UOC.
- 2.2. Chairperson and Secretary shall hold an informal meeting with new members to discuss responsibilities of members, ERC processes and procedures.
- 2.3. The members should attend training/workshops pertaining to the functions of the ERC regularly and maintain a training record in the member file.
- 2.4. New members shall be given the priority in the trainings
  - 2.4.1. Training should include training in Standard Operating Procedures, Research Ethics and human subject protection in compliance with FERCSL and other national and international guidelines.
- 2.5. It is the responsibility of members to obtain information and attend training courses, workshops and conferences that are announced periodically on websites, bulletin boards and the media.
- 2.6. Keeping the training records fill in the Training Record as per Annex 4 to record the training/workshop/conference activities in chronological order. A copy of the certificates of such training must be retained in the ERC office.



#### SOP - 006 Submission Procedure for New Applications Version 2 23/04/2018



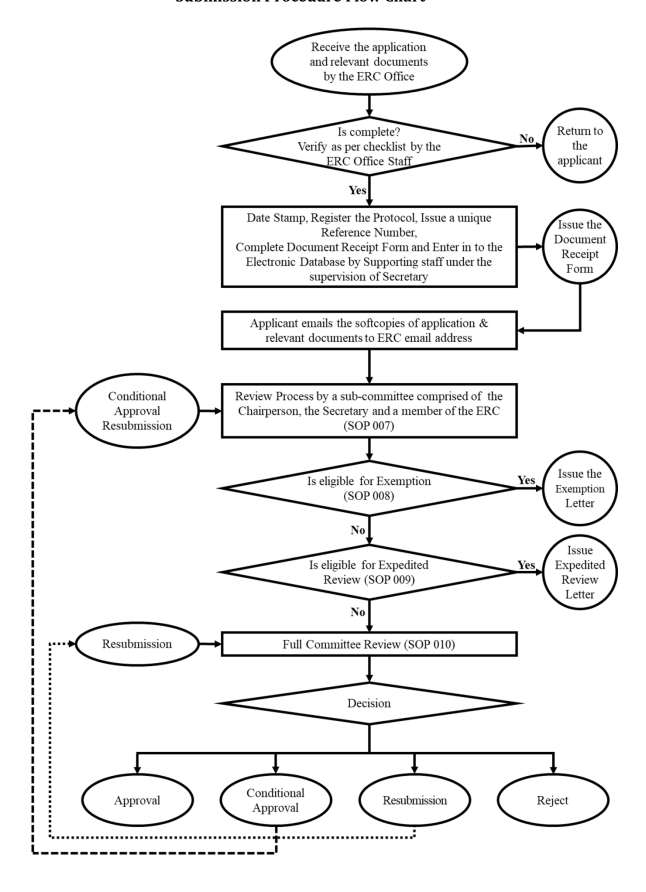
#### 1. Purpose: To describe the procedure for submission of new applications

New protocol submission includes initial submission of new protocols, resubmission of corrections/amendments and continuing review of approved protocols. It is the responsibility of the Secretary to receive, record, and distribute the protocols among the reviewers.

- 2.1. Applications must be submitted to the Secretary/ERC in the format prescribed by the ERC (Annex 05) and shall include all necessary documents. ERC application is available in the PGIM/UOC website.
- 2.2. Guidelines to fill the ERC applications are available in the PGIM/UOC website (http://pgim.cmb.ac.lk/?page\_id=1957).
- 2.3. Applications should be accompanied by the following documents:
  - a. Covering letter: covering letter should be signed by the applicant. If the applicant is a postgraduate trainee of the PGIM, covering letter should be submitted through the supervisor who is officially assigned to the applicant.
  - b. Declaration of Applicant (Annex 06)
  - c. Submission Check List. (Annex 07)
  - d. Research Protocol (03 copies).
  - e. Information Sheet (Annex 08) and Consent Form (Annex 09) in English, and in Sinhala and Tamil where appropriate (03 copies).
  - f. Other relevant documents (i.e. questionnaires) in English, and in Sinhala and Tamil where appropriate (03 copies).
  - g. Approval letter from the relevant Board of Study for postgraduate study protocols.
  - h. Updated Curriculum Vitae of the principal investigator and all the co-investigators as per Annex 10. In general, each CV should not be more than 2-3 pages, unless a complete CV is specifically requested for.

- i. Online payment receipt
- 2.4. Supporting staff in the ERC office ensures that all required forms and documents are submitted along with the application under the supervision of the Secretary.
- 2.5. Upon receipt of complete protocol, supporting staff of the ERC office should issue a unique registration number and enter the protocol into the electronic database. The format of the number should be ERC/PGIM/current year/serial number.
- 2.6. Document Receipt Form will be issued upon receipt of complete application along with all the necessary documents as per Annex 11.
- 2.7. A compressed/zipped folder containing soft copies of all the documents relevant to the application should be emailed to the ERC at <a href="erc@pgim.cmb.ac.lk">erc@pgim.cmb.ac.lk</a> within 24 hours of receipt of Document Receipt Form. The subject of the email should be ERC Registration Number followed by the last name of the applicant (eg. ERC/PGIM/2018/XXX Perera).
- 2.8. Upon receipt of an email from the principal investigator, all the protocols will be circulated among all ERC members via email.
- 2.9. A fee will be charged for applications as per Table 01.
- 2.10. Duly completed applications are accepted by the ERC office from Monday through Friday (except on public holidays) during office hours (9.00am to 4.00pm).
- 2.11. Deadline of applications for the regular monthly meeting shall be the close of business on the last working day of the previous month.
- 2.12. In the event of a public health emergency, such as the investigation of a disease outbreak or a disaster relief operation, the investigators may request a be reviewed expeditiously. In such proposal to instances. the Chairperson/Secretary may call emergency meeting the an subcommittee/full committee to discuss such protocols.

#### **Submission Procedure Flow Chart**





#### SOP - 007 Initial Review Process Version 2 23/04/2018



#### 1. Purpose: To describe the procedure for initial review of new protocols

The initial review includes initial screening of new protocols and deciding the review type. It is the responsibility of the subcommittee comprised of the Chairperson, the Secretary, and an assigned member to conduct the initial review. However, if any of the subcommittee members have a conflict of interest related to any proposal, another member from the ERC should be appointed for the subcommittee to review such proposals.

#### 2. Detailed Instruction:

2.1. The subcommittee meets weekly on Monday and screen all the new proposals received within the previous week. Subcommittee assesses the degree of risk involved and decides the review type.

#### 2.2. Degree of Risk

There are three levels of risk associated with human research as follows: No risk, Minimal risk, and More than Minimal Risk. The degree of risk involved in particular research will be determined based on these risk levels. This categorization should be applied as defined in the FERCSL guidelines.

#### 2.3. Types of review:

Based on the degree of risk, a proposal will be subjected to one of the review types

#### 2.3.1. Exemption from review

No risk is associated and proposals are exempted from ethics review when there is no possibility of harm arising as a result of the conduct of the research project or when the information being collected is available from the public domain.

#### 2.3.2. Expedited review

A proposal is considered for expedited review when the research procedures present no more than minimal harm to the research

participants or communities. In this case, the proposal is reviewed by the subcommittee.

#### 2.3.3. Full committee review

All research protocols with more than minimal risk to human subjects are reviewed by two ERC members as per SOP 10, using the prescribed format (Annex 05 Part B), who present the protocol to the ERC followed by a general discussion and a consensus decision. All the member of the ERC is expected to go through such proposal and provide their comments at the discussion.



#### SOP - 008 Exempted From Review Process Version 2 23/04/2018



#### 1. Purpose: To describe the procedure for new protocols exempted from review

Protocols which carry less than minimal risk fall under this category. It is the responsibility of the subcommittee comprised of the Chairperson, the Secretary, and an assigned member of the ERC to grant approval for exemption.

- 2.1. At the weekly subcommittee meeting, new proposals received within the previous week will be reviewed and the proposals with less than minimal risk will be exempted from review.
- 2.2. Proposals that fulfil any of the following conditions are exempted from review
  - a. Does not involve the collection or use of individual-level data or community level data on sensitive topics
  - b. All data to be used are freely available in the public domain
  - c. Research on cadavers and death certificates provided such research reveals no personally identifiable data
  - d. Audits or educational practices
- 2.3. Applicants whose applications qualify for an exemption will be informed by the Secretary.
- 2.4. Applications which are eligible for exemption from the review will be submitted to the next ERC meeting for ratification of the decision of the subcommittee.
- 2.5. A formal letter of exemption will be issued only after confirmation of the subcommittee's decision by the ERC (Annex 12).



#### SOP - 009 Expedited Review Process Version 2 23/04/2018



#### 1. Purpose: To describe the procedure for expedited review of new protocols

Protocols that carry a minimal risk to the participants or the community fall under this category. It is the responsibility of the subcommittee comprised of the Chairperson, the Secretary, and an assigned member of the ERC to grant approval.

- 2.1. The subcommittee meets weekly on Monday and screen all the new proposals received within the previous week. Subcommittee assesses the degree of risk involved and decides the review type.
- 2.2. The subcommittee may undertake an expedited review of proposals with minimal risk and those on non-sensitive topics under following circumstances
  - a. The participants are not considered a vulnerable group
  - b. The topic of research is not considered a sensitive topic
- 2.3. Applicants whose applications qualify for expedited review by the subcommittee will be informed by the Secretary of the results.
- 2.4. Applications which are eligible for expedited review will be submitted to the next ERC meeting for ratification of the decision of the subcommittee.
- 2.5. A formal letter of approval will be issued only after the confirmation of the subcommittee's decision by the ERC (Annex 13).



### SOP – 010 Full Committee Review Process

Version 2 23/04/2018



#### 1. Purpose: To describe the procedure for full board review of new protocols

Protocols which carry more than minimal risk fall under this category. Proposals which were not considered for exemption nor expedited review as needing full board review also will be reviewed.

- 2.1. The subcommittee meets weekly on Monday and screen all the new proposals received within the previous week. Subcommittee assesses the degree of risk involved and decides the review type. Proposals with more than minimal risk will undergo full board review.
- 2.2. If the proposal is not exempted nor undergone expedited review, then the subcommittee will assign primary reviewers based on their expertise. Two scientific reviewers and two non-scientific reviewers will be assigned to each protocol coming under this category.
- 2.3. The scientific reviewers are tasked to review technical soundness and related ethical issues while the non-scientific reviewers are tasked to review the informed consent process and forms.
- 2.4. The Secretary prepares the proposals for primary review and circulates among the assigned reviewers. Primary reviewers will review the protocols using review forms (SOP 005 Section B). Based on their preference hard copies will be circulated. Soft copies of all the proposals that undergo full board review shall be emailed to all the ERC members.
- 2.5. Decision making: When there is a quorum, the decision is arrived at by consensus. If consensus is not possible, voting is carried out. Only members who are present are allowed to participate in the voting. However, members who are joining the meeting through video conferencing and teleconferencing are also eligible to vote.

- 2.6. Opinions of absent members that are transmitted by mail or telephone or fax may be considered by the attending members during the discussion but may not be counted as votes or quorum for formally convened full board meetings.
- 2.7. External reviewer/s who are experts in the subject may be invited where necessary to offer their views, but external reviewer/s should not participate in the decision-making process. However, his/her comments must be recorded.
- 2.8. The full board review of a research proposal will result in one of the following actions
  - 2.8.1. Approved: The research proposal is approved as submitted. This does not preclude the Committee from sending comments for the consideration of the research team or requesting proof of approval by the local ERC/ERB/ IRB or proof of clinical trial registration in the Sri Lanka Clinical Trial Registry when appropriate.
  - 2.8.2. Conditional approval: If the full board approves a research proposal in principle subject to minor modifications ('Conditional Approval'), the revised project proposal submitted by the proponent will be reviewed and approved by the subcommittee. When the requirements are met, a letter of approval will be issued. Such revised proposals will not be taken up for the full board review. If required, submitted responses will be sent to primary reviewers by the subcommittee. If the primary reviewers are not satisfied with the response, the Secretary will request the applicant to provide further clarifications.
  - 2.8.3. Revise and resubmit: The research proposal is not approved as submitted either because there is insufficient information to make a decision or the proposal is not ethically sound. However, the proposal can be resubmitted for full board review after addressing all the comments of the first review. The revised documents will be discussed in a full board meeting.
  - 2.8.4. Reject: The research proposal is ethically or scientifically unacceptable.



#### SOP - 011 External/Independent Reviewers



Version 2 23/04/2018

### 1. Purpose: To describe the procedure for assigning external/independent reviewers

ERC will seek the advice of an external reviewer when the committee lacks the expertise among its members to review specific subject/technical areas. It is the responsibility of the subcommittee comprised of the Chairperson, the Secretary, and an assigned member to assign external/independent reviewers.

- 2.1. ERC maintains a list of external/independent reviewers who are experts in different subject areas.
- 2.2. The subcommittee may invite external/independent reviewers when they think the expertise within the ERC is not sufficient to evaluate a particular proposal.
- 2.3. ERC must be satisfied that such experts have no conflicts of interest in relation to the project under consideration arising from any personal involvement or participation in the project, any financial interest in the outcome or any involvement in competing research. Such person(s) shall be required to provide an undertaking of confidentiality and shall not be entitled to vote on any matter.



#### SOP - 012 Conflicts of Interest Version 2 23/04/2018



### 1. Purpose: To describe the procedure for reporting and handling conflicts of interest of the ERC members

The purpose of this SOP is to describe the procedure for reporting and handling of conflict of interest of the ERC members. It is the responsibility of all ERC members to understand, accept and declare any conflicts of interest before the ERC meeting.

- 2.1. An ERC member shall inform the Chairperson/Secretary if he/she has a conflict of interest, financial or otherwise, in a project or other related matter(s) to be considered by the ERC prior to the commencement of the meeting.
- 2.2. The ERC will determine if this results in a conflict of interest for the member and, if so, the member will withdraw from the meeting until the ERC's consideration of the relevant matter has been completed. The member shall not be permitted to adjudicate on the research.
- 2.3. All declarations of conflicts of interest and the resolutions of the same shall be minuted.



#### SOP - 013 Preparation of Agenda Version 2 23/04/2018



### 1. Purpose: To describe the procedure and format of the agenda for a meeting of the ERC

This SOP describes the process and format of the agenda. It is the responsibility of the Secretary to prepare and share the agenda.

- 2.1. The Secretary of the ERC will prepare an agenda for each meeting of the ERC.
- 2.2. All completed applications and relevant documents received by the ERC office by the agenda closing date will be included in the agenda.
- 2.3. The Secretary will circulate the agenda and associated documents among members of the ERC at least seven (07) calendar days prior to the respective meeting.
- 2.4. Agenda will include the following items
  - 1. Announcements/Welcome/Excuses
  - 2. Declaration of conflicts of interest by ERC members
  - 3. Confirmation of minutes of the previous meeting
  - 4. Matters arising from the previous minutes
  - 5. Training
  - 6. New applications
    - 6.1. Exempted New Protocols for Ratification
    - 6.2. New Protocols Subjected to Expedited Review for Ratification
    - 6.3. New Protocols For Full Board Review
  - 7. Previously considered unapproved applications for approval
  - 8. Amendments/extensions to approved protocols
  - 9. Progress/final reports of the approved protocols
  - 10. Reports of subcommittees
  - 11. Amendments to SOPs
  - 12. Correspondence

- 13. Any other business
- 14. Date, time and venue for the next meeting



#### SOP - 014 Conduct of Meetings Version 2 23/04/2018



#### 1. Purpose: To describe the procedure for the conduct of ERC meetings

This SOP describes the procedure for the conduct of the ERC meeting. It is the responsibility of the Chairperson and the Secretary to inform members and facilitate the conduct of regular and special meetings of the ERC.

- 2.1. The ERC shall meet on a monthly basis.
- 2.2. Members may attend ERC meetings in person or via teleconference or video conference. Members who are unable to attend a meeting should contribute prior to the meeting through written submissions to the Secretary of the ERC. The minutes should record the submission of written comments.
- 2.3. A quorum must be present in order for the ERC to reach a final decision on any agenda item. A quorum shall exist when more than one-third of the members are present including at least one member falling under category 2.3.1.d or 2.3.1.e described in SOP 002.
- 2.4. The Chairperson may cancel a scheduled meeting if a quorum cannot be achieved. Should this occur, the ERC will convene another meeting within fourteen (14) working days of the cancelled meeting to ensure all agenda items are taken up for discussion.
- 2.5. In exceptional circumstances, the Chairperson shall decide to proceed with the meeting even in the absence of a quorum. In such circumstances, decisions made by the ERC must be ratified by at least one member falling under category 2.3.1.d or 2.3.1.e described in SOP 002.
- 2.6. Meetings will not be restricted for an allocated time. Meetings will continue until all agenda items have been considered. All deliberations will be conducted in a manner that is non-offensive, unbiased, sensitive, and inclusive.
- 2.7. The ERC meeting will be conducted in private to ensure confidentiality and open discussion. Members will be advised of the venue in the meeting agenda.

- 2.8. Notwithstanding paragraph 2.7 above, the ERC may agree to the presence of visitors or observers at a meeting.
- 2.9. Any member of the ERC who has any interest, financial or otherwise in a proposal or other related matter(s) considered by the ERC, must declare such interests beforehand.



#### SOP - 015 Preparation of Minutes Version 2 23/04/2018



#### 1. Purpose: To describe the procedure and format of minutes of the ERC meeting

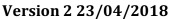
This SOP describes the administrative procedure for preparation ,review, approval and distribution of ERC meeting minutes. It is the responsibility of the Secretary to prepare the minutes and to ensure the quality and validity of the minutes after the meeting is over. It is the responsibility of the Chairperson to review and approve the minutes sent to him by the Secretary.

- 2.1. The Secretary of the ERC will prepare the minutes of each meeting of the ERC as per the template given in Annex 14.
- 2.2. All completed applications and relevant documents received by the ERC office by the agenda closing date will be included in the agenda.
- 2.3. The format of the minutes will include the following items:
  - 1. Attendance
  - 2. Announcements/Welcome/Excuses
  - 3. Declaration of Conflicts of interest
  - 4. Proceedings of the previous meeting
  - 5. Matters arising from the previous minutes
  - 6. Training
  - 7. New applications
    - 7.1. Exempted New Protocols for Ratification
    - 7.2. New Protocols Subjected to Expedited Review for Ratification
    - 7.3. New Protocols For Full Board Review
  - 8. Previously considered applications for approval
  - 9. Amendments/extensions of approved proposals
  - 10. Progress/Final reports of the approved proposals
  - 11. Reports from subcommittees
  - 12. Amendments to SOPs
  - 13. Correspondence

- 14. Any other matters
- 15. Next meeting and Close
- 2.4. The minutes should include the recording of decisions taken by the ERC as well as a summary of relevant discussions. This includes reference to views expressed in writing by absent members.
- 2.5. In relation to the review of new applications or amendments, the minutes shall record the ERC's decision and any requests for additional information, clarification or modification of the proposal.
- 2.6. In recording a decision made by the ERC, any significant dissenting view or concern will be noted in the minutes.
- 2.7. To encourage free and open discussion and to emphasize the collegiate character of ERC deliberations, particular views shall not be attributed to particular individuals in the minutes, except in circumstances where a member seeks to have his/her opinions or objections recorded.
- 2.8. Declarations of conflicts of interest by any member of the ERC and the absence of the member concerned during the ERC deliberation of the relevant application will be minuted (Refer to SOP 012 regarding Conflicts of Interest).
- 2.9. The minutes will be produced as soon as possible following the relevant meeting and will be checked by the Chairperson for accuracy.
- 2.10. The minutes will be circulated among all members of the ERC as an agenda item for the next meeting. All members will be given the opportunity to seek amendments to the minutes prior to their ratification. The minutes will be formally ratified at the next ERC meeting.
- 2.11. The confirmed and amended minutes of each meeting (with the inclusion of revisions if any) will be filed in the 'Minutes File'.
- 2.12. The extracts of minutes shall be sent to the Director, PGIM/UOC and Board of Management, PGIM/UOC. The extracts will consist of the titles of the approved proposals and the decision of the ERC that would need Board of Management ratification.



### SOP – 016 Notification of Decisions of ERC





### 1. Purpose: To describe the procedure for the notification of ERC decisions concerning review of new applicants

The purpose of this SOP is to ensure proper completion, distribution and filing of communications with investigators. It is the responsibility of all ERC members, including the Secretary and the Chairperson, to complete a written communication record for telephone or interpersonal discussions related to past, present and/or future studies and/or processes involving the ERC.

- 2.1. The Secretary of the ERC will prepare the ERC Decision letter two weeks after the monthly ERC meeting.
- 2.2. Decision letters can be collected from the ERC office two weeks after the monthly meeting
- 2.3. If the ERC determines that further information, clarification or modification is required for the consideration of a project, the correspondence to the principal investigator should clearly articulate the reasons for this determination, and clearly set out the information that is required (Annex 15). Where possible, requests for additional information/ clarification/ modification should refer to the FERCSL Guidelines or other relevant documents including legislation.
- 2.4. The ERC shall endeavour to openly communicate with applicants to resolve outstanding requests for further information, clarification or modification of protocols relating to ethical issues. The ERC may nominate one of its members to communicate directly with the applicant or invite the applicant to attend the relevant ERC meeting.
- 2.5. Notification of ethical approval will be in writing and will contain the following information. A standard letter will be issued, in the format set out in Annex 16.
  - 1. Title of the project
  - 2. Name of the principal investigator(s)

- 3. Unique ERC identification number
- 4. Version number and date of all documentation reviewed and approved by the ERC including protocols, information sheets, consent forms questionnaires etc.
- 5. Date of the ERC meeting at which the project was first considered
- 6. Date of the ERC's approval
- 7. Conditions of the ERC's approval, if any
- 8. Duration of the ERC's approval
- 2.6. The research project may not commence until written notification of ethical approval is received and non-adherence to this requirement amounts to ethical misconduct.
- 2.7. Any extensions for ethics approval for conducting the research project should be requested before the expiry of the validity indicated in the previous ethics clearance approval.
- 2.8. If the ERC determines that a project is ethically unacceptable, the notification of the ERC"s decision will include the grounds for rejecting the project with reference to the FERCSL Guidelines or other relevant pieces of legislation. A standard rejection letter will be issued, in the format set out in Annex 17.
- 2.9. The status of the project shall be updated on the ERC's register of received and reviewed applications.



#### SOP – 017 Amendments and Extensions to Approved Protocols Version 2 23/04/2018



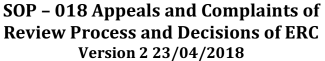
### 1. Purpose: To describe the procedure for the submission and ERC review of requests for amendments and extensions to approved protocols.

This SOP applies to proposals submitted to the ERC, PGIM/UOC undergoing amendments or subsequent extensions after initial approval. It is the responsibility of the Secretary to forward such requests to the ERC considering the need for expedited review or full committee review in consultation with the Chairperson.

- 2.1. Approval for proposed changes to approved research protocols or to the conduct of the research, including extensions to the length of ERC approval, must be sought by the principal investigator in writing.
- 2.2. Requests shall outline the nature of the proposed changes and/or request for an extension, reason/s for the request, and an assessment of any ethical implications arising from the request on the conduct of the research. All amended documents must have the changes highlighted. The request for extension must be accompanied by a current progress report of the study.
- 2.3. Expedited review of requests for minor amendments and extensions may be undertaken by the ERC subcommittee between scheduled meetings at the discretion of the Chairperson or the Secretary and in accordance with SOP 009, on the condition that it is ratified at the next ERC meeting.
- 2.4. Where an urgent protocol amendment is required for safety reasons, the Chairperson may review and approve the request. In such circumstances, the ERC will review the decision at its next meeting.
- 2.5. All other requests for amendments shall be reviewed by the ERC at its next meeting, provided the request has been received by the ERC office by the agenda closing date.
- 2.6. The ERC will report in writing to the principal investigator, advising of the ethical approval of the proposed amendment and/or request for extension and

- that the amended research may commence, within seven (7) working days of the meeting at which the request was considered (this may be the full ERC meeting or the subcommittee meeting).
- 2.7. If the ERC determines that further information, clarification or modification is required for the consideration of the request for amendment or extension, the correspondence to the investigator should clearly articulate the reasons for this determination, and clearly set out the information that is required. Where possible, requests for additional information/clarification/modification should refer to the relevant pieces of legislation
- 2.8. All reviewed and approved requests for amendments and extensions shall be recorded in the relevant protocol specific file and, where appropriate, in the ERC"s register of received and reviewed applications.







## 1. Purpose: To describe the procedure for receiving and handling appeals or complaints regarding the ERC's review process and decisions

This SOP applies to complaints/appeals submitted to the ERC, PGIM/UOC by applicants who are not satisfied with the ERC review/decision. It is the responsibility of the Chairperson to investigate such complaints/appeals.

- 2.1. An applicant who is not satisfied with the outcome of the ERC's decision may complain to the Chairperson detailing in writing the grounds of the concern or complaint. Complaints may also be made to the Director, PGIM/UOC.
- 2.2. The Chairperson will inform the Director, PGIM/UOC as soon as possible of any complaints received by him/her. The Director, PGIM/UOC will inform the Chairperson as soon as possible of any complaints received by him/her. The Director, PGIM/UOC will send a letter of acknowledgement to the complainant, outlining the following mechanism.
- 2.3. Appeal/complaint will be tabled at the next ERC meeting and the Chairperson will appoint a panel of three members, excluding the members who originally reviewed the protocol, to investigate the appeal/complaint.
- 2.4. The decision of the panel will be discussed at the subsequent ERC meeting. The decision of the ERC will be informed to the applicant within three months of the complaint.
- 2.5. If the complainant is not satisfied with the outcome of the Chairperson's investigation, then he/she can refer the complaint to the Director, PGIM/UOC or request that the Chairperson do so.
- 2.6. The ERC will provide the Director, PGIM/UOC with all relevant information about the complaint/concern, including:
  - 1. The appeal.
  - 2. Material reviewed.

- 3. The decision of the review panel.
- 4. Any other relevant documentation.
- 2.7. Further action if necessary will be at the discretion of the Director, PGIM/UOC, and may include one or more of the following:
  - 1. Agree with the ERC decision.
  - 2. Appoint a three-member panel comprising the Director, PGIM/UOC or his/her nominee and two senior academics nominated by the Board of Management of the PGIM/UOC to review and give recommendations.
  - 3. Any action deemed suitable considering the recommendations of the panel.



### SOP – 019 Monitoring of Approved Research Projects Version 2 23/04/2018



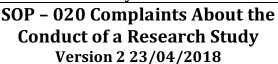
1. Purpose: To describe the procedure for monitoring research projects approved by the ERC to ensure compliance with ethical approval.

The ERC will monitor approved protocols to ensure compliance with its ethical approval. The purpose of this SOP is to describe the procedure for monitoring research protocols approved by the ERC to ensure compliance with ethics approval.

- 2.1. The ERC shall monitor approved projects to ensure compliance with the conditions for ethical approval. In doing so, it may request and discuss information on any relevant aspects of the project with the investigators at any time. In particular, the ERC shall require the investigators to provide annual progress reports (Annex 18) and a final report (Annex 19) at the completion of the study.
- 2.2. The ERC shall require the following information in the progress reports
  - 1. Progress to date or outcome in the case of completed research
  - 2. Maintenance and security of records
  - 3. Compliance with the approved protocol
  - 4. Compliance with conditions of approval
  - 5. Changes related to study investigators and sources of funding
- 2.3. The ERC may undertake random site visits as part of monitoring.
- 2.4. The ERC shall require, as a condition of approval of each project, that investigators immediately report anything which might warrant a review of the ethical approval of the proposal, including, but not limited to:
  - any unforeseen events that might affect the continued ethical acceptability of the project Standard Operating Procedures – ERC, PGIM/UOC
  - 2. new information from other published or unpublished studies which may have an impact on the continued ethical acceptability of the project, or which may indicate the need for amendments to the protocol.

- 2.5. The ERC shall require, as a condition of approval of each project, that investigators inform the ERC, giving reasons if the research project is discontinued before the expected date of completion.
- 2.6. Where the ERC is of the opinion that the research project is not being conducted in accordance with the approved protocol, the ERC may withdraw approval. In such circumstances, the ERC shall inform the principal investigator and the institution as well as any Regulatory Authority of such withdrawal of approval in writing, and recommend to the institution that the research project be discontinued, suspended, or that other necessary steps be taken.







1. Purpose: To describe the procedure for receiving, handling and responding to complaints concerning the conduct of a project approved by the ERC.

The ERC shall receive complaints from research participants, researchers, or other interested individuals regarding the conduct of approved research projects. The contact details of the ERC must be included in the Patient/ Participant Information Sheet for each project.

- 2.1. Any concern or complaint received will be forwarded to the Chairperson of the ERC. The Chairperson is responsible for obtaining a written complaint stating the grounds of the concern. Upon receiving this, the ERC will be notified as soon as possible.
- 2.2. The ERC shall send a letter of acknowledgement to the complainant and a letter of notification to the principal investigator outlining the complaint and the mechanism for investigating (described below) the complaint.
- 2.3. Where the complaint concerns a serious matter within the jurisdiction of the Ministry of Health or other institution, the Chairperson shall consider referral of the complaint to the Ministry of Health or the relevant governing body.
- 2.4. A panel consisting of a minimum of three (03) members will be appointed by the ERC to conduct an investigation of the complaint. This panel upon completion of the investigation shall make recommendations to the ERC on the appropriate course of action. Based on the seriousness of the violation one or more of the following action will be recommended.
  - 1. Amendments to the protocol
  - 2. Warning and increased monitoring by the ERC
  - 3. Suspension of the project
  - 4. Termination of the project
  - 5. Other appropriate action to resolve the complaint

- 2.5. Such action will be taken within three months of receiving a written complaint.
- 2.6. The complainant shall be informed of the action taken. He/she will also be informed of his /her right to refer the complaint to the Director, PGIM/UOC if he/she is not satisfied with the decision of the ERC.





### SOP – 021 Dealing with Protocol Deviations/Violations Version 2 23/04/2018

## 1. Purpose: To describe the procedure for the process for reporting and handling of protocol deviations and violations

The purpose of this SOP is to describe how the ERC, PGIM/UOC provides instructions for taking action and maintaining records, when investigators fail to follow the procedures written in the approved protocol or fail to comply with national/ international guidelines for the conduct of human research, including those who fail to respond to the ERC, PGIM/UOC requests.

- 2.1. The ERC shall require, as a condition of approval of each proposal, that researchers report to the ERC of any protocol deviation or violation as soon as possible but no later than one (01) calendar month of its first knowledge.
- 2.2. The report should include,
  - 1. ERC reference number
  - 2. Details of the site
  - 3. Details of protocol deviation/violation
  - Reason(s) for deviation patient related/investigator related/other (specify)
  - 5. Details of the reporter Name, address, telephone number, other administrative information
  - 6. Measures taken by the investigators to deal with the violation and to avoid future occurrences
- 2.3. All reported deviations and violations will be dealt with by a subcommittee consisting of Chairperson, Secretary and an assigned ERC member and will be informed to the ERC, at the next meeting



### SOP - 022 Record Keeping Version 1.0 23/04/2018



### 1. Purpose: To describe the procedure for the preparation and maintenance of records of the ERC activities

It is the responsibility of the Secretary ERC to prepare and maintain written/electronic records of all the ERC activities. The supporting staff of the ERC will prepare and maintain a confidential electronic and/or paper record for each application received and reviewed

- 2.1. Supporting staff of the ERC will prepare and maintain a confidential electronic and/or paper record for each application received and reviewed and shall record the following information under the supervision of the Secretary:
  - 1. ERC unique identification number
  - 2. Title of the project
  - 3. The principal investigator(s) with contact details
  - 4. Name of the responsible institution or organization
  - 5. Date of Submission
  - 6. Date of review at an ERC meeting and the decision(s) taken at this meeting
  - 7. Decision/s of the ERC approval or non-approval with date/s
  - 8. Approval or non-approval of any changes to the project
  - 9. Terms and conditions, if any, of approval of the project and
  - 10. Type of approval, whether approval was by expedited review.
- 2.2. The paper file shall contain a hard copy of the application, including signatures, and any relevant correspondence including that between the applicant and the ERC, all approved documents and other material used to inform potential research participants.

- 2.3. All relevant records of the ERC, including applications, membership, minutes, correspondence, and progress/final reports will be kept as confidential files.
- 2.4. To ensure confidentiality, all documents provided to ERC members, which are no longer required, are to be disposed of in a secure manner, such as shredding.
- 2.5. All records pertaining to research protocols shall be held for sufficient time to allow for future reference. The minimum period for retention will be five (5) years. Files which are no longer required for retention shall be electronically archived.
- 2.6. A register of all the applications received and reviewed shall be maintained in accordance with the FERCSL and other national/international guidelines.



### SOP – 023 Review of Standard Operating Procedures



Version 1.0 23/04/2018

1. Purpose: To describe the procedure for the process for reviewing and amending SOPs within the ERC, PGIM/UOC.

It is the responsibility of the Secretary of the ERC to appoint the SOP subcommittee to amend the SOPs by following the same procedures, format and coding system when drafting or editing any SOP of the ERC.

- 2.1. The Terms of reference and Standard Operating Procedures shall be reviewed at least every three years and amended as necessary.
- 2.2. SOPs may be amended at any time if a need arises for such amendments
- 2.3. The SOPs may be amended by following the procedure below
  - 1. Any member of the ERC can propose an amendment to the SOPs in writing.
  - 2. The proposed amendment shall be submitted in writing to the Secretary to be placed in the agenda of the next available Ethics Review Committee meeting for consideration and possible adoption by at least two-thirds of the committee members present and voting. Any member unable to attend such a meeting may register their views in writing
  - 3. The Chairperson shall send the amendment to the Director, PGIM/UOC for review and approval, if appropriate.
  - 4. The amendment shall come into effect once approved by the Director and Board of Management, PGIM/UOC.

### References

- Declaration of Helsinki (DoH) of the World Medical Association (WMA), 2013.
- Ethics review committee guidelines, Forum of Ethics Review Committees, Sri Lanka 2007.
- Ethical review committee Guidelines. Sri Lanka Medical Association. Colombo 1999.
- International Conference on Harmonization, Guidance on Good Clinical Practice (ICH GCP) E6 (R1) 1996.
- International Ethical Guidelines for Epidemiological Studies Prepared by the Council for International Organizations of Medical Sciences (CIOMS) 2008.
- National Ethical Guidelines for Health Research in Nepal. Nepal Health Research Council, Nepal 2001.
- Standards and Operational Guidance for Ethics Review of Health-Related Research with Human Participants, World Health Organization 2011.
- Standards Operating Procedures, Ethics Review Committee, Faculty of Medical Sciences, University of Sri Jayewardenepura 2012.
- Standard Operating Procedures, Ethics Review Committee, Faculty of Medicine, University of Colombo 2016.
- Standards Operating Procedures, Ethics Review Committee, Faculty of Medicine, University of Kelaniya 2018.
- WMA Declaration of Taipei on Ethical Considerations regarding Health Databases and Biobanks. Available at:
  - http://www.wma.net/en/30publications/10policies/d1/index.html [Accessed 12 November 2016]
- The Nuremberg Code. Reprinted from Trials of War Criminals before the Nuremberg Military Tribunals under Control Council Law No. 10, Vol. 2. Washington, D.C., U.S. Government Printing Office, 1949. Available at
- WHO Operational Guidelines for Ethical Review Committee that review biomedical research, Geneva 2000. Available at <a href="https://www.who.int/tdr/publications/">www.who.int/tdr/publications/</a> [Accessed on 25th August 2015]







Category	Fee
1. Trainers and Staff of the PGIM of UOC	Free
2. Trainees of the PGIM of UOC	Rs. 2000
3. Extended Faculty of the PGIM of UOC	Rs. 5000
4. Industry-sponsored projects	Rs. 7500

- Where the application falls under two or more of these categories, whichever the highest fee will apply.
- All the payments should be made through PGIM online payment system prior to the submission and a hard copy of the receipt should be submitted along with the application.
- Fees once paid will not be refunded.







Th	is agreem	ent is	s made a	and $\epsilon$	entered int	o on this				da	ay of
			by and	d bet	ween Ethio	cs Review Com	ımittee, Po	stgr	adua	te Inst	itute
of	Medicine	, Ur	niversity	of	Colombo	(hereinafter	referred	to	as	ERC)	and
								(ho	older	of	NIC
nu	mber			)	of		(herein	afte	r ref	erred	to as
the	e "member	") W	HEREAS	the	member h	as agreed to se	erve on the	e afo	resai	id ERC	, and
in	which cap	acity	the men	ıber	will have a	ccess to Confid	dential Info	orma	tion	in the	ERC;
AN	D WHERE	AS th	ie memb	er ha	s acknowl	edged and agre	eed that the	e con	nmit	tee has	and
sha	all continu	e to	have sol	e rig	hts to Conf	idential Inform	nation and	l has	agr	eed to	hold
the	same in s	strict	confide	nce o	during and	after the men	iber's peri	od o	f ser	vice w	ithin
the	e ERC.										

And it is hereby agreed as follows

### 1. Interpretation

"Confidential information" shall include all information of a confidential and proprietary nature provided or made available to the member by the ERC including but not limited to the research proposals and documents, techniques, intellectual property and processes and such other information related to the ERC but shall not include information which is or becomes publicly available other than through the faults of the member.

### 2. Obligations of the member

a. To maintain the highest degree of secrecy and keep as confidential any Confidential Information which the member may be granted access to, or which may be available to, or which member receives on behalf of the ERC or in the capacity of the member of the ERC by any means and to use such Confidential Information only in duty authorized manner in the interest of the ERC and for the purpose of fulfilling functions and responsibilities arising as a member of the ERC.

- b. Not at any time during or after service within the ERC, for any reason, disclose or permit to be disclosed any Confidential Information to any third party or to use such Confidential Information for personal use without the express prior written approval of the ERC.
- c. On termination of the period of membership within the ERC, for whatever reason to the ERC all property, documents and papers in the member's possessions or control relating to the inter alia of the ERC.
- d. That in the event of a break of any of the conditions mentioned above, the ERC shall be entitled to injunctive relief and/or specific performance to enforce the conditions set out above.

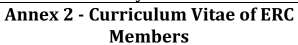
### 3. The legal compulsion to disclose

In the event that the member becomes legally compelled to disclose any Confidential Information, the member shall give prompt notice in writing of such facts to the ERC so that ERC has an opportunity to seek a protective order or another remedy. In the event that such protective order or other appropriated remedy is not sought by the ERC or is sought but is not obtained, the member will nevertheless disclose only that portion of the Confidential Information as is necessary to comply with its obligations under law and shall use reasonable endeavours to obtain an appropriate court order or other reliable assurance that Confidential treatments will be accorded to Confidential Information so disclosed.

4. The member hereby unconditionally accepts and acknowledges that having regard to the nature of the ERC and the functions and duties of the member of the ERC the member considers the terms and conditions imposed herein has been fair and reasonable.

Signature of the Member	Date
Signature of the Chairperson of the ERC	Date







Personal Information	n		
Name	Rev/Prof/Dr/Mr/M	S	
Current Designation			
Home Address			
Contact Number			
Email address			
<b>Educational Qualific</b>	ations		
Bachelor's degree			
Postgraduate			
degrees			
Work Experience			
Employment	Designation	Workplace	Period
Present			
Previous 1			
Previous 2			
Previous 3			
Training in Ethics			
Training 1			
Training 2			
Training 3			
Publications			
Date	-	Signature	of the ERC Member



### **Annex 3 - Letter of Appointment**



<Reference No.>

<Date>

<Name of the member>

<Address of the member>

Dear < Name of the member>,

## Appointment as a Member of the Ethics Review Committee, Postgraduate Institute of Medicine, University of Colombo

I am pleased to inform you that you have been appointed as a member of the Ethics Review Committee of the Postgraduate Institute of Medicine, by the Board of Management for a period of three years effective from <Date of appointment>.

The Secretary, ERC, PGIM/UOC will provide you with the Standard Operating Procedures (SOPs) of the ERC/FGS with which you are expected to be familiar. You are required to sign a confidentiality agreement on the assumption of duties.

The PGIM will indemnify you in respect of liabilities that may arise in the course of bona fide conduct of duties as an ERC member.

Your contributions as a member of the ethics review committee will be greatly appreciated.

Yours sincerely,

<Signature of the Director>
<Name of the Director>
Director, PGIM





### Annex 4 - Training Record

Name of the ERC Member: Rev/Prof/Dr/Ms/Mr								
Name of the Training	Training Provider	Venue	Date					



### **Annex 5 - Application Form for Scientific and Ethical Review**



requests
requests
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) (

1. Title of the Research Project: Enter title of the research project here

### 2. Details of the Investigators:

Title, Name, Designation and Affiliation	Role	Signature
	Principal	
	Investigator	

3.	<b>Contact Details</b>	of the	<b>Principal</b>	Investigator:
•		01 0110	III OI POI	. III OUUL MUUL

If Yes, Details:

	3.1 Postal Address	Enter the name of Principal Investigator
	3.2 Email Address	Enter the name of Principal Investigator
	3.3 Telephone	Enter the name of Principal Investigator
ŀ.	Nature of the study: Observational/non-inte Research database/info	
5.		tial date of enrolment of participants) and ending ellection) dates (retrospective approval will not be given tarted)  End Date:
j.		d of Study/Specialty Board approved the research
	Yes: No: If Yes, Board of Study/SDetails:	] pecialty Board:
		r this study been requested earlier from ERC, ERC? (if you have received ethics approval already, please roval)
3.	Funding (if any) Name and Address of th Amount:	e funding source:
9.	Do you believe the pro	oposed project has conflicts of interest?
	Yes: No:	

### Part B - Protocol Check List

Under each category, indicate the protocol section of the research proposal. If a particular category is not relevant to your study, indicate it as 'N/A'

	Scientific validity	Protocol		valuation		
		page/s	Ac	cepta	ble	Comments
			Yes	No	N/A	
1	Title					
2	Research problem					
3	Research questions/ hypothesis					
4	Objectives					
5	Study setting					
6	Study design					
7	Study population (giving inclusion-					
	exclusion criteria)					
8	Sample size					
9	Sampling method					
10	Measurements / variables					
11	Study instruments					
12	Procedures to ensure the quality of					
	data					
13	Plan for analysis					
14	Ethical considerations					
15	Budget (if relevant)					
16	Work plan and time frame					
17	Justification for a replication study, if					
	your study is a replication					

	Social Value	Protocol	Reviewer Evaluation				
		page/s	Acceptable			Comments	
			Yes	No	N/A		
1	Benefits of the study to the						
	community/society						
2	Plan for dissemination of study						
	findings						
3	Scientific importance of the study						

	Risk Benefit Assessment	Protocol	Reviewer Evaluation						
		page/s	Acceptable			Comments			
			Yes	No	N/A				
1	Potential risks to the participants								

2	Potential benefits to the participants			
3	Justification for risks against benefits			
4	Steps taken to minimize risks			
5	Support provided to participants			
	(medical, educational, other)			

	Participants rights and consent	Protocol		Reviewer Evaluation			
		pages	Ac	cepta	ble	Comments	
			Yes	No	N/A		
1	Procedure for recruiting the						
	participants						
2	Information provided to the						
	participants						
3	Procedure for requesting informed						
	consent						
4	Procedure for requesting proxy						
	consent						
5	Procedure for requesting assent						
	(subjects between 12y to 18y)						
6	Procedure for withdrawing consent						
7	Incentives provided to participants						
8	Procedure for participants to ask						
	questions / register complaints						
9	Participants right to decline						
	consent without losing entitled						
	benefits						

	Confidentiality and Privacy	Protocol		Reviewer Evaluation				
		page/s	Ac	cepta	ble	Comments		
			Yes	No	N/A			
1	Steps to ensure confidentiality of							
	data							
2	Justification for collecting personal							
	identification data							
3	Steps taken to ensure privacy							
	during data collection							
4	How long data and samples will be							
	kept							
5	Who will have access to the data							
6	Procedure for storage of data and							
	samples							
7	Procedure for disposal of data							

	Fair participant selection and	Protocol	<b>Reviewer Evaluation</b>				
	vulnerability	page/s	Acceptable		!	Comments	
			Yes	No	N/A		
1	Justification for selection of the						
	study population						
2	Justification for conducting the						
	study in a vulnerable population						

	Responsibilities of the	Protocol	Reviewer Evaluation			
	researcher	page/s	Acceptable		)	Comments
			Yes	No	N/A	
1	Ethical, legal, financial issues					
	related to the study					
2	Any conflicts of interest and how					
	the researcher plans to manage					
	them					
3	Permissions from relevant					
	institutions / authorities					
4	Collaborations with the relevant					
	stakeholder					
5	Provision of medical /					
	psychological care to the					
	participants					
6	Qualifications of the research team					
	to handle the research study	_				

	Foreign funded studies	Protocol	Reviewer Evaluation				
		page/s	Acce	ptable	!	Comments	
			Yes	No	N/A		
1	Justification for conducting the						
	study in SL						
2	Relevance of the study to SL						
3	Post research benefits to SL						
4	The sharing of intellectual property						
	rights						
5	How the results will be conveyed to						
	authorities in SL						

	Information Sheet / Consent	Section	Reviewer Evaluation				
	Form	in Info.	Acce	ptable	)	Comments	
		sheet	Yes	No	N/A		
		consent					
		form					
1	Purpose of the study						
2	Voluntary participation						
3	Duration of the study and						
	responsibilities of the participants						
4	Potential benefits						
5	Risks, Hazards, Discomforts						
6	Incentives / Reimbursements						
7	Confidentiality						
8	Contact person for the participants						
9	Understanding of information						
	provided by the researcher						
10	Agreement of the participant to						
	provide information / samples						
11	Consent for dissemination of						
	research findings	_					
12	Appropriate translation of the						
	information sheet						
13	Appropriate translation of the						
	consent form						

-				
Decision of the reviewer	r:			
Approved				
Conditional approval				
Approve with revisions				
Reject				
Comments of the Review	ver:			
Name of the Reviewer:				
Signature of the Reviewer	·	 	Date:	



### **Annex 6 - Applicant Declaration**

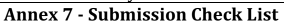


### **Declaration**

As the principal investigator of this project, my signature confirms that I will ensure that all procedures performed under the project will be conducted in accordance with all relevant national and international policies and regulations that govern research involving humans and cadavers. I understand that if there is any significant deviation from the project as originally approved, I must submit an amendment to the ERC for approval prior to its implementation. I have submitted all significant previous decisions by this or any other ERC and /or regulatory authorities relevant to the proposed study. I declare that I am not seeking approval for a study that has already been commenced or has already been completed.

Signature of principal investigator	Date:
Full name of principal investigator:	







For Office Use Only: Application Number: PGIM/ERC/20/	Date Received://20
To be Filled by the Applicant:	
Title: Name of the Applicant: Rev/Prof/Dr/Mr/Ms	

Document	Version	Date
1. Application form (3 copies)		
2. Detailed research proposal (3 copies)		
3. All study instruments (questionnaires/interview		
guides/checklist/data extraction forms)		
English (3 copies)		
4. Study instruments - Sinhala (if applicable) (3 copies)		
5. Study instruments - Tamil (if applicable) (3 copies)		
6. Information sheet - English (3 copies)		
7. Information sheet - Sinhala (if applicable) (3 copies)		
8. Information sheet - Tamil (if applicable) (3 copies)		
9. Consent forms - English (3 copies)		
10. Consent forms - Sinhala (if applicable) (3 copies)		
11. Consent forms - Tamil (if applicable) (3 copies)		
12. Any other relevant documents - English (3 copies)		
13. Any other relevant documents - Sinhala (3 copies)		
14. Any other relevant documents - Tamil (3 copies)		
15. Curriculum vitae of all investigators (1 copy)		
16. Approval letter from relevant Board of Study/Specialty		
Board for postgraduate studies (1 copy)		
17. Online payment receipt (1 copy)		



### **Annex 8 - Sample Information Sheet**



### <Title of the project>

I/We <name of principal investigator/s>, a <Designation> attached to <institute/s of affiliation> would like to invite you to take part in a research project titled <Non-technical Title> conducted by <Names of Investigators> at <Study Site>

### 1. Purpose

The objective/s of the study in non-technical terms

### 2. Voluntary participation

Your participation in this study is voluntary. You are free to not participate at all or to withdraw from the study at any time despite consenting to take part earlier. There will be no loss of medical care or any other available treatment for your illness or condition to which you are otherwise entitled. If you decide not to participate or withdraw from the study you may do so at any time.

### 3. Duration, procedures of the study and participant's responsibilities

This study will be conducted over a period of < anticipated duration of study>. If you volunteer to participate in this study, we will ask you to do the following:

- a. We will ask you to <take part /visit the clinic> for < duration of each visit and number of visits> over the course of a total of about < expected duration of participation>.
- b. You will need to <the procedure/s of the research including what happens at each visit in simple terms and how the participant has to take part in the study>

### 4. Potential benefits

Participation in this study may benefit you/others by < all the actual and potential benefits>

### 5. Risks, hazards and discomforts

<Any potential or actual risks, hazards and discomforts should be clearly defined>

6. Reimbursements

You would be paid a sum of Rs. < if any payment to the participant, indicating the amount,

when it would be paid and any conditions attached to it> OR you will not be paid any

sum of money for participating in this study

7. Termination of study participation

You may stop participating in this study at any time (with no penalty or effect on

medical care or loss of benefits). Please notify the investigator as soon as you decide

to withdraw your consent.

8. Confidentiality

Confidentiality of all records is guaranteed and no information by which you can be

identified will be released or published. These data will never be used in such a way

that you could be identified in any way in any public presentation or publication

without your express permission.

9. Clarifications

If you have questions about any of the tests / procedures or information, please feel

free to ask any of the persons listed below.

<The names and contact information of investigator/s>

<postal address, email address, telephone numbers>

If you have any clarification, concerns, or complaints related to this research project,

you may contact the Ethics Review Committee, Postgraduate Institute of Medicine,

University of Colombo.

ERC Office Address: Ethics Review Committee, Postgraduate Institute of Medicine,

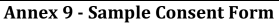
University of Colombo, 160, Prof. Nandadasa Kodagoda Mawatha, Colombo 07.

Telephone: 0112-689266 (between 9am and 4pm on working days)

Email: erc@pgim.cmb.ac.lk

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### <Title of the Research Project>

To be completed by the participant (Please tick the appropriate b	ox)	
Have you read the information sheet? (Please keep a copy for yourself)	Yes	No
2. Have you had an opportunity to discuss this study and ask any questions?		
<ul><li>3. Have you had satisfactory answers to all your questions?</li><li>4. Have you received enough information about the study?</li><li>5. Do you understand that you are free to withdraw from the study at any time, without having to give a reason and without affecting your future medical care?</li></ul>		
6. Sections of your medical notes, including those held by the investigators relating to your participation in this study may be examined by other research assistants. All personal details will be treated as strictly Confidential. Do you give your permission for these individuals to have access to your records?		
7. Have you had sufficient time to come to your decision? 8. Do you agree to take part in this study?		
Who explained to you about the		
study:		
Signature of the participant:Full name:	Date:	
1 UII IIUIIIC		
To be completed by the investigator/person obtaining consent I have explained the study to the above participant and he/she has in willingness to take part in this study.	dicated	her
Signature of Investigator:	Date:	

Full name:

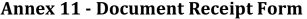




## Annex 10 - Curriculum Vitae of the Applicants

Personal Information	n			
Name	Rev/Prof/Dr/Ms/M	<b>I</b> r		
Current Designation				
Home Address				
Contact Number				
Email address				
Educational/Profess	ional Qualifications			
Bachelor's degree				
Postgraduate				
degrees				
Work Experience				
Employment	Designation	Workplac	e	Period
Present				
Previous 1				
Previous 2				
Publications (list up	to 5 most relevant t	o the proposed	d study)	
Ongoing Research Pr	rojects (other than t	his project)		
	_	<u> </u>		
Date		5	Signature	of the Applicant







Please insert the **Title** and the **Name of the Applicant** Title: **Name of the Applicant**: Rev/Prof/Dr/Mr/Ms For Office Use Only: This checklist will be filled and signed by the person who receives the application at ERC, PGIM/UOC office. Application Number: PGIM/ERC/20\_\_/\_\_\_ Date Received: \_\_\_/\_\_/20\_\_\_ 1. Covering Letter 2. Submission Check List 3. Declaration of Applicant 4. Application form (3 copies) 5. Detailed research proposal (3 copies) 6. All study instruments in English (questionnaires/interview guides/checklist/data extraction forms) with Sinhala and Tamil translations where relevant (3 copies) 7. Information sheet in English with Sinhala and Tamil translations where relevant (3 copies) 8. Consent forms in English with Sinhala and Tamil translations where relevant (3 copies) 9. Any other relevant documents in English with Sinhala and Tamil translations where relevant (3 copies) 10. Curriculum vitae of all investigators 11. Approval letter from the relevant Board of Study (if applicable) 12. Online payment receipt **Instructions for the applicants:** Please email all the documents within 24 hours of receipt of this checklist to erc@pgim.cmb.ac.lk as a compressed/zipped folder. The subject line should be the Application Number followed by the last name/surname of the applicant (Eg PGIM/ERC/2017/xxx Perera). Received by: ..... ..... ...... Name of the Staff Member Signature Date



### **Annex 12 - Exemption Letter**



<ERC Reference Number> <Date>

<Name of the Principal Investigator> <Address of the Principal Investigator>

Dear < Name of the Principal Investigator >,

### <Title of the Proposal>

Investigators:<Names of the investigators>

Thank you for submitting the above research proposal to the ERC of the Postgraduate Institute of Medicine. I am pleased to inform you that the study was exempted from the ethics review by the ERC at its meeting held on <meeting date> after reviewing the following documents submitted by you.

Document	Version No	Submission Date
Protocol	<version></version>	<date></date>
Information sheet	<version></version>	<date></date>
Consent form	<version></version>	<date></date>
Study Instrument	<version></version>	<date></date>

Thank you. Yours sincerely,



### **Annex 13 - Expedited Approval Letter**



<ERC Reference Number>
<Date>

<Name of the Principal Investigator><Address of the Principal Investigator>

Dear < Name of the Principal Investigator>,

### <Title of the Proposal>

Investigators:<Names of the investigators>

Thank you for submitting the above research proposal to the ERC of the Postgraduate Institute of Medicine. I am pleased to inform you that the study was approved by the ERC at its meeting held on <meeting date> after an expedited review of the following documents submitted by you.

Document	Version No	Submission Date
Protocol	<version></version>	<date></date>
Information sheet	<version></version>	<date></date>
Consent form	<version></version>	<date></date>
Study Instrument	<version></version>	<date></date>

The approval is valid until one year from <approval date>. You may submit a written request for renewal/extension of the approval, along with a progress report. Please note that you are required to inform the ERC about the following:

- Any unanticipated events involving potential risks to study subjects
- Any deviations in protocol
- Any changes to the documents listed above

You are required to submit the final report to the ERC/PGIM with the following declaration:

"the research was conducted in accordance with the proposal for which approval was granted by the ERC of PGIM" within three (03) months upon the completion of the study.

Thank you. Yours sincerely,







### Minutes of the <Meeting No> Meeting of the PGIM Ethics Review Committee <Date> at <Time> at <Venue>

### -CONFIDENTIAL-

### Attendance

Name	Position	<year></year>								
		Date Mon	Date Mon	Date Mon	Date Mon	Date Mon	Date Mon	Date Mon	Date Mon	Date Mon
<name chairperson="" of="" the=""></name>	Chairperson									
<name of="" secretary="" the=""></name>	Secretary									
<name member="" of="" the=""></name>	Member									
<name member="" of="" the=""></name>	Member									
<name member="" of="" the=""></name>	Member									
<name member="" of="" the=""></name>	Member									
<name member="" of="" the=""></name>	Member									
<name member="" of="" the=""></name>	Member									
<name member="" of="" the=""></name>	Member									
<name member="" of="" the=""></name>	Member									
<name member="" of="" the=""></name>	Member									

**P** – Present **E** – Excused **V**- Virtually Participated **A** – Absent **L** – on Leave

The <Meeting No.>th ERC meeting was called to order at <time>. <Name of the person who chaired the meeting> chaired the meeting.

### <Meeting No.>.1 Announcements/Welcome/Excuses

<announcements, welcome and excuses if any>

### <Meeting No.>.2 Declaration of Conflict of Interest

<Member> - <ERC Ref No of the protocol>

### < Meeting No. > .3 Confirmation of the Minutes

Minutes of the <last Meeting No>th ERC meeting held on <last meeting date> were confirmed. Proposed by <name of the member who proposed > and seconded by <name of the member who seconded>

### <Meeting No.>.4 Matters arising from the Minutes

<matters arising from the minutes of the last meeting>

### <Meeting No.>.5 Training

### <Meeting No.>.5.1 In-house Training

<in-house training topic, name of the
member who conducted the training>

### <Meeting No.>.5.2 Other trainings

<details if any>

<Meeting No.>.6 New Protocols

<Meeting No.>.6.1 Exempted New Protocols for Ratification

<Meeting No.>.6.1.1 <ERC Ref No.><Title><Name of the Principal Investigator>

### <Meeting No.>.6.1.2 New Protocols Subjected to Expedited Review for Ratification

<Meeting No.>.6.1.3 <ERC Ref No.><Title><Name of the Principal Investigator>

### <Meeting No.>.6.2 New Protocols For Full Board Review

<Meeting No.>.6.2.1 <ERC Ref No.><Title><Name of the Principal Investigator>

ERC No:	<erc no.="" ref=""></erc>		Date Subm	Date Submitted:			<submission date=""></submission>		
Applicant	<name a<="" of="" td="" the=""><td>pplicant</td><td>&gt;</td><td></td><td></td><td></td><td></td><td></td><td></td></name>	pplicant	>						
Study Design	<type of="" study=""></type>								
Documents	Application	n Protocol		Instrument		IS/ICF			
				Е	S	T	Е	S	T
Version	<version no=""></version>	<version no=""></version>		<version no=""></version>			<version no=""></version>		
Internal	<name 2nd<="" <name="" of="" td="" the=""><td colspan="5"><name of="" person="" reviews="" sinhala<="" td="" the="" who=""></name></td></name>			<name of="" person="" reviews="" sinhala<="" td="" the="" who=""></name>					
reviewers	1st Primary	Primary	7	version>					
	Reviewer> Reviewer>		<name of="" person="" reviews="" tami<="" td="" the="" who=""><td>iews Tamil</td></name>				iews Tamil		
	ver			versi	on>				
ERC Discussion	<discussion points=""></discussion>								
Recommendation	<erc approval="" i.e.="" of="" recommendation.="" type=""></erc>								
Remarks	<details any="" if=""></details>								

### <Meeting No.>.7 Previously Considered Protocols

ERC No:	<erc no.="" ref=""></erc>		Date Subm	e Submitted:			<submission date=""></submission>		
Applicant	<name a<="" of="" td="" the=""><td>pplicant</td><td>&gt;</td><td></td><td></td><td></td><td></td><td></td><td></td></name>	pplicant	>						
Study Design	<type of="" study=""></type>								
Documents	Application Protocol		otocol	Ins	strum	ent		IS/ICI	F
				Е	S	T	Е	S	T
Version	<version no=""></version>	<version no=""></version>		<version no=""></version>		<version no=""></version>			
Internal	<name of="" td="" the<=""><td><name< td=""><td>of the <math>2^{nd}</math></td><td colspan="4"><name of="" person="" reviews="" sinhala<="" td="" the="" who=""></name></td></name<></td></name>	<name< td=""><td>of the <math>2^{nd}</math></td><td colspan="4"><name of="" person="" reviews="" sinhala<="" td="" the="" who=""></name></td></name<>	of the $2^{nd}$	<name of="" person="" reviews="" sinhala<="" td="" the="" who=""></name>					
reviewers	1st Primary Primary		7	version>					
	Reviewer>	viewer> Reviewer>		<name of="" person="" reviews="" tam<="" td="" the="" who=""><td>iews Tamil</td></name>			iews Tamil		
			versi	on>					
ERC Discussion	<discussion points=""></discussion>								
Recommendation	<erc approval="" i.e.="" of="" recommendation.="" type=""></erc>								
Remarks	<details any="" if=""></details>	·						·	

### <Meeting No.> .8 Amendments/Extensions to Approved Protocols

<meeting No.>.8.1 <ERC Ref No.> - <Title>

### <Meeting No.>.9 Progress/Final Reports of approved proposals

<meeting No.>.9.1 <ERC Ref No.> - <Title>

### < Meeting No.>.10 Reports of the Subcommittees

<Details, if any>

### < Meeting No.>.11 Amendments to SOPs

<Details, if any>

### <Meeting No.>.12 Correspondence

<Details, if any>

### <Meeting No.>.13 Any Other Business

<Details, if any>

### <Meeting No.>.14 Date, Time and Venue for the next meeting

<Date> <Time> <Venue> for the next meeting

Meeting adjourned at <adjourned time>

<Signature of the Secretary>

<Name of the Secretary>

Secretary, ERC, PGIM/UOC



# **Annex 15 - Resubmission Letter**



<ERC Reference Number> <Date>

<Name of the Principal Investigator> <Address of the Principal Investigator>

Dear < Name of the Principal Investigator >,

### <Title of the Proposal>

Investigators:<Names of the investigators>

Thank you for submitting the above research proposal. ERC at its meeting held on <meeting date> reviewed the following documents submitted by you.

Document	Version No	Submission Date
Protocol	<version></version>	<date></date>
Information sheet	<version></version>	<date></date>
Consent form-	<version></version>	<date></date>
Study Instrument	<version></version>	<date></date>

The ERC has suggested following modifications prior to the consideration for approval.

<suggested modifications>

Please resubmit the revised proposal with a covering letter, within three (03) months of this letter. Please underline all the changes in the proposal and indicate the changes in the covering letter in a table with four columns indicating reviewers comments, original wording, the responses/revisions and page numbers.

Thank you. Yours sincerely,



### Annex 16 - Approval Letter



<ERC Reference Number>
<Date>

<Name of the Principal Investigator><Address of the Principal Investigator>

Dear < Name of the Principal Investigator >,

### <Title of the Proposal>

Investigators:<Names of the investigators>

Thank you for submitting the above research proposal. I am pleased to inform you that the study was approved by the ERC at its meeting held on <meeting date> after reviewing following documents submitted by you.

Document	Version No	Submission Date
Protocol	<version></version>	<date></date>
Information sheet	<version></version>	<date></date>
Consent form-	<version></version>	<date></date>
Study Instrument	<version></version>	<date></date>

The approval is valid until one year from the meeting date stated above. You may make a written request for renewal/extension of the validity, along with the submission of a progress report.

Please note that you are required to inform the ERC about the following:

- Any unanticipated events involving potential risks to study subjects
- Any deviations in protocol
- Any changes to the documents listed above

You are required to submit the final report to the ERC/PGIM with the following declaration:

"the research was conducted in accordance with the proposal for which approval was granted by the ERC of PGIM" within three (03) months upon the completion of the study.

Thank you. Yours sincerely,



### **Annex 17 - Rejection Letter**



<ERC Reference Number> <Date>

<Name of the Principal Investigator> <Address of the Principal Investigator>

Dear < Name of the Principal Investigator >,

### <Title of the Proposal>

Investigators:<Names of the investigators>

Thank you for submitting the above research proposal. I regret to inform you that your study was NOT approved by the ERC at its meeting held on <meeting date> based on following documents submitted by you.

Document	Version No	Submission Date
Protocol	<version></version>	<date></date>
Information sheet	<version></version>	<date></date>
Consent form-	<version></version>	<date></date>
Study Instrument	<version></version>	<date></date>

Main reason/s for this decision is/are <reasons for rejection>

You may submit a new application after addressing all the above comments.

Thank you. Yours sincerely,



### Annex 18 - Progress Report



ERC Ref No.	
Title	
Details of Principal Investigator	
Name of the Principal	
Investigator	
Address of the Principal	
Investigator	
Phone Number	
Details of the Study	
Date of Approval	
Study Start Date	
Progress	
Progress to the date	
Maintenance and security	
of records	
Compliance with the	
approved protocol	
Protocol	
deviations/violations	
Presentations/publications	
related to data gathered in	
this study	
Any other	
·	
Date	Signature of the PI

This report should be submitted every six (06) months



### Annex 19 - Final Report



ERC Ref No.	
Title	
Details of Principal Investigator	
Name of the Principal	
Investigator	
Address of the Principal	
Investigator	
Phone Number	
Details of the Study	
Date of Approval	
Study Start Date	
Study End Date	
Number of Participants	
Main Findings	
Protocol Deviations/Violations	
Presentations/publications	
Any other	
Date	Signature of the PI