

To
 The Chairperson
 Institutional Ethics Committee
 Symbiosis International (Deemed University)
 Lavale, Pune 412115
 Email id: iec@siu.edu.in

Application for Review of Research Protocol / Project by Ethics Committee
 (All the fields are mandatory. Mention 'NA' if item not applicable.)

A. Basic Information

1	Research Study Title	
2	Project ID / PRN (In case of PhD Scholar)	
3	Full Name of Principal Investigator (PI)	
4	Designation of PI	
5	Faculty / Department	
6	Affiliation of PI	
7	Name sponsor for the Research, if any	

B. Details about Co-Investigators

Sr. No.	Full Name of Co-Investigator	Designation	Affiliation
1.			
2.			
3.			
4.			
5.			
6.			

C. Research Related Information

1	Type of Review requested	exemption/expedited/full review
2	Are you seeking waiver of consent ? if Yes, Specify Reason	
3	Whether study approved by IRC / RAC/ Any other scientific committee (Attach copy of approval)	Yes / No

4	Whether study needs collection of data from the participants/constituents of SIU? [If Yes, attach the approval from the Data Access Review Committee (DARC) of SIU]	Yes / No
5	Does your project involve ethical review by other ethics committee (Attach copy)	Yes / No
6	Type of Study	Drug Trial or Clinical Trial
		Biomedical and Health Research
		Multi-centric / Single Center
		Any other (specify)
7	If Clinical Trial	Does study involve Drugs, Devices or Vaccines (specify)
8		Does study involve a change in dose, route or indication of the drug? (if yes specify)
9		Statutory authority's permission is obtained? (if yes date of permission)
10		CTRI Registration Number (if registered)
11		Phase of the clinical study
12		Is similar study being conducted elsewhere? (if yes, attach details)

D. Checklist of the documents to be attached with this form

		Yes	NA
1	Protocol / Plan of study		
2	Case Report Form / Study Tool / Questionnaire		
3	Subject Information Document: In English		
4	Subject Information Document: In Local Language		
5	Approval by RAC/IRC or equivalent Technical Committee		
6	Informed Consent Form: In English		
7	Informed Consent Form: In Local Language		
8	Investigator's Brochure (Only Applicable to Drug Trials)		
9	Subject Recruitment Materials: In English		
10	Subject Recruitment Materials: In Local Language		
11	Details of remuneration to the participants		
12	Details of the research grant		
13	CVs of PI and Co-Investigators		

E. Participant Related Information

	Does your study involve any of the following	Yes	No
1	Research involving pregnant women or the human fetus		
2	Participants unable to give consent due to high dependency on medical care		
3	Participants with a cognitive impairment or mental illness		
4	Participants involved in illegal activities		
5	Use of invasive Interventions / Therapies		
6	Human Genetics / stem cells related research		
7	Projects involving ionizing radiations		

8	Projects involving active concealment or planned deception of participants		
9	Collection of identifiable personal information, without permission from the person identified.		
10	Risk of harm to participants (More than discomfort)		
11	Participants unable to give consent due to language difficulties		
12	Are there any risk to the researchers (e.g. unsafe environment, trouble spots)		
13	Are there any other risks not covered in the above assessment that you consider maybe relevant		

F. If your answer is ‘yes’ to any items on the checklist E, you may comment any additional information related to Ethical consideration in following text box.

G. Vulnerable Populations

	Does the research specifically target participants from any of the following groups?	Yes	No
1	Children under age of 18 years		
2	Participants with physical disability		
3	Participants whose ability to give consent is impaired		
4	Residents of custodian institutions		
5	Participants with dependent relationship with the researchers (e.g. teacher-student, doctor-patient, professional-client)		
6	Research involving sensitive cultural issues		

H. If your answer is ‘yes’ to any items on the checklist G, you may comment any additional information related to Ethical consideration in following text box.

I. Elements of an informed consent document

The consent form can be created with the help of National Ethical Guidelines for Biomedical and Health Research Involving Human Participants, ICMR (2017) and New Drugs and Clinical Trials Rules (2019).

As per ICMR guidelines (2017), an informed consent form must include the following:

1. Statement mentioning that it is research
2. Purpose and methods of the research in simple language
3. Expected duration of the participation and frequency of contact with estimated number of participants to be enrolled, types of data collection and methods
4. Benefits to the participant, community or others that might reasonably be expected as an outcome of research
5. Any foreseeable risks, discomfort or inconvenience to the participant resulting from participation in the study
6. Extent to which confidentiality of records could be maintained, such as the limits to which the researcher would be able to safeguard confidentiality and the anticipated consequences of breach of confidentiality
7. Payment/reimbursement for participation and incidental expenses depending on the type of study
8. Free treatment and/or compensation of participants for research-related injury and/or harm
9. Freedom of the individual to participate and/or withdraw from research at any time without penalty or loss of benefits to which the participant would otherwise be entitled
10. The identity of the research team and contact persons with addresses and phone numbers (for example, PI/Co PI for queries related to the research and Chairperson/Member Secretary/ or helpline for appeal against violations of ethical principles and human rights)

J. Key Terms:

1. **Deception in human subject research** means deliberately misleading subjects about the nature of a study.
2. **Concealment** means deliberately withholding certain information. Studies involving deception or concealment must meet all criteria for a waiver or alteration of informed consent.
3. **IRC / RAC approval** means technical approval of the study at the level of institute or the University (in case of Ph D students) respectively.
4. **Trouble Spots** means areas where researcher might find it unsafe due to happenings in the field for example areas prevalent with highly infectious diseases, disaster prone areas.

K. Declarations:

1. I/we declare that the information filled in this form is complete and correct.
2. I/we confirm that the study will be conducted in accordance with all applicable statutory guidelines and regulations.

Signature of the Principal Investigator

Date: