Application Form for Initial Review All India Institute of Medical Sciences, Madurai EC Ref. No.(for office use):								
General Instructions: a) Tick one or more as applicable. Mark NA if not applicable b) Attach additional sheets if required								
SEC	TION A - BASIC	INFORMATION						
<ul> <li><b>ADMINISTRATIVE DETAILS</b> <ul> <li>(a) Name of Organization:</li> <li>(b) Name of the Ethics Committee:</li> <li>(c) Name of Principal Investigator:</li> <li>(d) Department/Division:</li> <li>(e) Date of Submission: click here to enter a date.</li> </ul> </li> <li>(f) Type of review requested<sup>1</sup>:</li> </ul>								
Exemption from Review	Expedited Revie	w 🔲 Full Committee Review 🗖						
Acronym/ Short title, (If any):								
<ul><li>(h) Protocol number(If any):</li><li>(i) Details of Investigators:</li></ul>		Version number:						
Name Designation and Qualification	Department and Institution	Address for communication <sup>2</sup>						
Principal Investigator/Guide								
Co-investigator/student/fellow								
(j) Number of studies where applicar		·						
i) Principal Investigator at tim	e of submission:	ii) Co-Investigator at time of submission:						
(k) Duration of the study:								
2. FUNDING DETAILS AND BUDGET								
<sup>1</sup> Refer to National Ethical Guidelines for Biomed review	ical & Health Research Inv	olving Human Participants 2017on Page 36 Table 4.2. for the types of						

<sup>2</sup>Include telephone/mobile, fax numbers and email id

	At site	In	te: India	Globally		
)	Self-funding		Institutional funding 🔲		unding agency	
		SECTIO	N B - RESEARCH RELA	TED INF	ORMATION	
. 0\	VERVIEW OF RE	ESEARCH				
(a)	Lay Summary	y of study <sup>3</sup> (wi	thin 300 words)			
(b)	Type of study	·				
	Basic Sciences Retrospective	السل	Clinical Epidemiological/ Public		Cross Sectional Case Control	
			Health			
	Prospective Qualitative		Socio-behavioural		Cohort Systematic Review	
	Quantitative		Biological		,	
	Mixed Metho	d 🗖	samples/Data			
			Any others (Specify)			
. M (a)	ETHODOLOGY Sample size/ At site		pants ( <i>as applicable)</i> India Globally			
	Sample size/ At site Control group	In Study G for the sample	India Globally	n case of q	ualitative study, menti	on the criteria
(a) (b)	Sample size/ At site Control group Justification f used for satu	In Study G for the sample iration xternal labora	India Globally Globally e size chosen ( <i>100 words</i> ); I tory/ outsourcing involved			
(a)	Sample size/ At site Control group Justification f used for satu	In Study G for the sample iration sternal labora	India Globally Group e size chosen ( <i>100 words</i> ); I			
(a) (b)	Sample size/ At site Control group Justification f used for satu Is there an ex How was the	In Study G for the sample iration sternal labora	India Globally Group e size chosen ( <i>100 words</i> ); I tory/ outsourcing involved lity of the study assessed?	for investig	gations? <sup>4</sup> Yes 🔲 No	
(a) (b)	Sample size/ At site Control group Justification f used for satu Is there an ex How was the Independen	In Study G for the sample iration sternal labora scientific qua nt external hin multi-	India Globally Group e size chosen ( <i>100 words</i> ); I tory/ outsourcing involved lity of the study assessed? Review by	for investig	gations? <sup>4</sup> Yes 🗖 No Review within	
(a) (b)	Sample size/ At site Control group Justification f used for satu Is there an ex How was the Independen review Review with	In Study G for the sample iration xternal labora e scientific qua nt external hin multi- arch group	India Globally Group e size chosen ( <i>100 words</i> ); I tory/ outsourcing involved lity of the study assessed? Review by Sponsor/Funder	for investig	gations? <sup>4</sup> Yes 🔲 No Review within Pl's institution	
(a) (b)	Sample size/ At site Control group Justification f used for satu Is there an ex How was the Independen review Review with centre resea Date of revie	In Study G for the sample iration xternal labora e scientific qua nt external hin multi- arch group	India Globally Group e size chosen ( <i>100 words</i> ); I tory/ outsourcing involved lity of the study assessed? Review by Sponsor/Funder	for investig	gations? <sup>4</sup> Yes 🔲 No Review within Pl's institution	NA NA N
(a) (b)	Sample size/ At site Control group Justification f used for satu Is there an ex How was the Independen review Review with centre resea Date of revie	In Study G for the sample iration xternal labora e scientific qua at external hin multi- arch group ew: of Scientific Co	India Globally Globally Group e size chosen (100 words); I tory/ outsourcing involved lity of the study assessed? Review by Sponsor/Funder No Review	for investig	gations? <sup>4</sup> Yes <b>No</b> Review within PI's institution Click here to	NA NA N

<sup>3</sup>Summarize in the simplest possible way such that a person with no prior knowledge of the subject can easily understand it. <sup>4</sup>If participant samples are sent outside for investigations, provide details of the same and attach relevant documentation such as an MTA/ MoU etc.

5. RE	ECRUITMENT AND RESEARCH PARTICIPANTS	
(a)	Healthy 🔲 Patient 🔲 Vul	nerable person/  Others Contended of the second sec
	Who will do the recruitment? Participant recruitment methods used:	
	Posters/ TV/Radio leaflets/Letters ads/Social media/Institution website	Patients /
	Others(Specify)	
(b)	<ul><li>i. Will there be vulnerable person/special groups i</li><li>ii. If yes, type of vulnerable person /special groups</li></ul>	
	Children under 18 yrs	Pregnant or lactating women
	Differently abled (Mental/Physical)	Employees/Students/Nurses/
	Elderly	Institutionalized
	Economically and socially disadvantaged Terminally III (stigmatized or rare diseases)	Refugees/Migrants/Homeless
	Any other (Specify):	
	iii. Provide justification for inclusion/exclusion	
	iv. Are there any additional safeguards to protect re	esearch participants?
(c)	Is there any reimbursement to the participant? If yes, Monetary 🔲 Non-monetary 🔲 Provide de	Yes 🗖 No 🗖
(d)	Are there any incentives to the participant?	Yes 🗖 No 🗖
	If yes, Monetary 🗖 Non-monetary 🗖 Provide det	ails
(e)	Are there any participant recruitment fees/ incentives for	or the study provided to the PI/ Institution?
	lf yes, Monetary 🗖 Non-monetary 🗖 Provide deta	ails Yes No
6. BEI	ENEFITS AND RISKS	

(a)	i. Are there any anticipated physical/social/	psychological discomforts/ risk to participants? Yes 🔲 No 🗖
	If yes, categorize the level of risk <sup>5</sup> : Less than Minimal risk	Minimal risk
	Minor increase over minimal risk or Low Risk ii. Describe the risk management strategy:	More than Minimal Risk or High Risk
(b)	What are the potential benefits from the stud	y? Yes No If yes, Direct Indirect
	For the participant	
	For the society/community	
	For improvement in science Please describe how the benefits justify the ri	sks
(c)	Are Adverse Events expected in the study <sup>6</sup> ?	Yes 🔲 No 🔲 NA 🗖
	Are reporting procedures and management st If Yes, Specify	rategies described in the study? Yes 🔲 No 🔲
7. II	NFORMED CONSENT	
(a)	Are you seeking waiver of consent? If yes, ple	ase specify reasons and skip to question 8. Yes 🔲 No 🗖
(b)	Version number and date of Participant Inform Version number and date of Informed Consent	
(c)	Type of consent planned for :	
	Signed consent  Verbal/ oral  consent	Witnessed Audio-Video consent (A/V) consent
	Consent from LAR For children<7 yrs (If so, specify from whom) consent	<ul> <li>Verbal assent</li> <li>from minor (7-</li> <li>from Minor (13-</li> <li>yrs) along</li> <li>with parental</li> <li>consent</li> <li>Written Assent</li> <li>Written Assent</li> <li>from Minor (13-</li> <li>18 yrs) along with</li> <li>parental consent</li> </ul>
(4)	Other <i>(specify)</i> <b>U</b> Who will obtain the informed consent?	
(d)	PI/Co-I Nurse/Counselor	Research Staff Other(Specify)
	Any tools to be used	
2.	1	edical & Health Research Involving Human Participants 2017. Page 6 in Table
• <i>T</i>	The term adverse events in this regard encompass both serious an	d non-serious adverse events. Version 2.0 04

(e)	Participant Information Sheet(PIS) and Informed Consent Form (ICF) English Local language other ( <i>specify</i> ) List the languages in which translations were done								
(f)	If translation has not been done, please justify Provide details of Consent requirement for previously stored samples if used in the study <sup>7</sup>								
(g)	Elements contained in the Participant Information Sheet(PIS) and Informed Consent Form (ICF)								
	Simple language 🔲 Data/ Sample 🔲 Compensation for study related injury 🔲								
	Risks and		sharing Need to recontact		Statement t	hat consent is voluntary			
	discomforts Alternatives to		Confidentiality		Commercial	ization/benefit sharing			
	participation Right to withdraw		Storage of		Statement t	hat study involves research			
	Benefits		samples return of research results		Use of photo	ographs/ identifying data			
	Purpose and procedure Others( <i>Specify</i> )		Payment for participation		Contact info Secretary of	rmation of PI and Member EC			
<b>8. P</b> (a	AYMENT/COMPENS ) Who will bear the PI	costs	related to participation		procedures <sup>8</sup> ? onsor	Other agencies(specify)			
(b	) Is there a provisio	n for fi	ree treatment of rese	arch re	elated injuries	? Yes 🗖 No 🕻			
(c	•	•	vide the treatment? ompensation of resea	arch re	lated SAE? If y	ves, specify. Yes 🗖 No			
	Sponsor 🗖 Ins	stitutio	n/ Corpus funds	F	Project grants	Insurance			
(d			r medical treatment o s during the study per		-	he relatedness is determin Yes 🔲 No			
(e	) Is there a provisior specify.	) for an	cillary care for unrela	ated illi	ness during th	e study period? If yes, pleas Yes 🔲 No	se NA		
-	ormation on re-consent requ Participants 2017,Page 54 lose undertaking from PI cor	in Sectio	n 5.8	ical Guide	elines for Biomedicc	al & Health Research Involving Humar	1		

<b>9. ST(</b> (a)	DRAGE AND CONFIDENTIALITY Identifying Information: Study Involves samples/data. If Yes, Specify	Yes 🔲 No 💭 NA 💭
	Anonymous/unidentified 🔲 Anonymized: Irreversibly reversibly coded 🗖 coded 🗖	Identifiable
	If identifiers must be retained, what additional precautions will be taken to / data is safeguarded? (e.g. data stored in a cabinet, password protected con	
(b)	Who will be maintaining the data pertaining to the study?	
(c)	Where will the data be analyzed <sup>9</sup> and by whom?	
(d)	For how long will the data be stored?	
(e)	Do you propose to use stored samples/data in future studies? If yes, explain how you might use stored material/data in the future?	Yes 🔲 No 🗖 Maybe 🗖
	SECTION D: OTHER ISSUES	
10. PU	BLICATION, BENEFIT SHARING AND IPR ISSUES	
(a)	Will the results of the study be reported and disseminated? If yes, specify.	Yes 🗖 No 🗖 NA 🗖
(b)	Will you inform participants about the results of the study?	Yes 🔲 No 💭 NA 💭
(c)	Are there any arrangements for continued provision of the intervention for once the study has finished? If yes describe in brief ( <i>Max 50 words</i> )	r participants, if effective, Yes 🔲 No 🔲 NA 🗖
(d)	Is there any plan for post research benefit sharing with participants? If yes,	specify Yes 🔲 No 🔲 NA 🗖
(e)	Is there is any commercial value or a plan to patent/IPR issues. If yes, Pleas	e provide details Yes 🔲 No 💭 NA 💭
(f)	Do you have any additional information to add in support of theapplication elsewhere in the form? If yes, provide the details.	n, which is not included Yes 🔲 No 🗖
<sup>9</sup> For	example, a data entry room, a protected computer etc.	

## SECTION E: DECLARATION AND CHECKLIST<sup>10</sup>

11. D	11. DECLARATION (Please tick as applicable)							
	I/We certify that the information	I/We certify that the information provided in this application is complete and correct.						
	I/We confirm that all investigators have approved the submitted version of proposal/related documents.							
		I be conducted in accordance with the latest ICMR National and Health Research involving Human Participants and other nes including responsible.						
		be conducted in accordance with the Drugs and Cosmetics Act nded from time to time, GCP guidelines and other applicable						
	I/We will comply with all policie institutions where this study will b	es and guidelines of the institute and affiliated/collaborating be conducted.						
	I/We will ensure that personnel will adhere to the provisions of th	performing this study are qualified, appropriately trained and e EC approved protocol.						
		in case of injury related to the study will be taken care of.						
	If applicable, I/We confirm that an provided, if applicable.	n undertaking of what will be done with the leftover samples is						
	I/We confirm that we shall submi	I/We confirm that we shall submit any protocol amendments, adverse events report, significant deviations from protocols, progress reports (if required) and a final report and also participate in						
	I/We confirm that we will maintai	n accurate and complete records of all aspects of the study.						
	I/We will protect the privacy of p and biological samples.	participants and assure safety and confidentiality of study data						
		the investigators, researchers and/or close relative(s), have no -Financial) with the sponsor(s) and outcome of study.						
	I/We have the following conflict o	f interest (PI/Co-PI):						
	1. 2.							
	I/We declare/confirm that all requirements wherever applicable	necessary government approvals will be obtained as per e.						
	Name of PI: Signature:	Click here to enter a date.						
	Name of Co-PI: Signature:	Click here to enter a date.						

12. C	Name of Guide: Signature: Click he Name of HOD: Signature: Click her				Enclosure	
S.No	Items	Yes	No	NA	No.	EC Remarks(lf applicable)
ADM	NISTRATIVE REQUIREMENTS	·		·		
1.	Cover letter					
2.	Brief CV of all Investigators					
3.	Good Clinical Practice (GCP) training of investigators in last 3 years					
4.	Approval of Scientific Committee					
5.	EC clearance of other centers*					
6.	Agreement between collaborating partners*					
7.	MTA between collaborating partners*					
8.	Insurance policy/certificate					
9.	Evidence of external laboratory credentials in case of an externally outsourced laboratory study QA/QC certification					
10.	Copy of contract or agreement signed with the sponsor or donor agency					
11.	Provide all significant previous decisions (e.g. those leading to a negative decision or modified protocol) by other ECs/Regulatory authorities for proposed study (whether in same location or elsewhere) and modification(s) to protocol					
PROP	OSAL RELATED					
12.	Copy of the detailed protocol <sup>11</sup>					
13.	Investigators Brochure (If applicable for drug/biologicals/device trials)					
14.	Participant Information Sheet(PIS) and Informed					

	Consent Form (ICF)(English a	nd translate	ed)							
15.	Assent form for minors (12-18 years) (English and Translated)									
16.	Proforma/Questionnaire / Interview guides/ Guides for (FGDs) (English and translate									
17.	Advertisement/material to posters etc)			fliers,						
PERM	IISSION FROM GOVERNING AU	JTHORITIES	T	I		1		1		
	Other Registration/ permissions	Required	Not required	Rece	ived	Appli dd/m	ed m/yy	EC Remark	S	
18.	CTRI					Enter				
19.	DCGI					Enter date				
20.	HMSC					Enter date				
21.	NAC-SCRT						date			
22.	ICSCR					Enter	date			
23.	RCGM					Enter	date			
24.	GEAC					Enter	date			
25.	BARC					Enter	date			
26.	Tribal Board						date			
27.	Others (Specify)						date			
ANY	OTHER RELEVANT INFORMATI	ON/DOCUM	IENTS RELA	TED T	O THE	STUDY	,			
	ltem		YES	NO	NA	Enclo no.	sure	EC remarks		
28.										
29.										

<sup>10</sup>These formats are adaptable and can be modified by the Ethics Committee members depending on their needs and requirements Acknowledgement for Receipt of Application (Copy to be provided to PI)

\*For multicentric research. MTA-Material transfer agreement; CTRI-Clinical Trial Registry-India; DCGI-Drug Controller General of India;HMSC-Health Ministry's Screening Committee;NAC-SCRT- National Apex Committee for Stem Cell Research and Therapy;IC-SCR-Institutional committee for Stem Cell Research;RCGM- Review Committee on Genetic Manipulation;GEAC- Genetic Engineering Approval Committee;BARC- Bhabha Atomic Research Centre

<sup>11</sup>Refer to National Ethical Guidelines for Biomedical and Health Research Involving Human Participants 2017, section 4 page no. 35Box 4.4(b)