Logo of the Institute

Application Form for Initial Review

(Name of the Institution)	EC Ref. No. (For office use):

General Instructions: a) Tick one or more options as applicable. Mark NA if not applicable b) Attach additional sheets if required

SECTION A - BASIC INFORMATION

ADMINISTRATIVE DETAIL	LS		
(a) Name of Organization	າ:		
(b) Name of Ethics Comr	nittee:		
(c) Name of Principal Inv	estigator:		
(d) Department/Division	:	(e) Date of subm	ission: dd mm yy
(f) Type of review reques	ited¹:		
Exemption from review	ew 🗆 Expedited rev	riew □ Full cor	mmittee review \square
(g) Title of the study:			
Acronym/ Short title	e, (If any): ny):		number:
Acronym/ Short title	e, (If any): ny):		
Acronym/ Short title	e, (If any):ny):rs: Designation and	Version Department and	
Acronym/ Short title (h) Protocol number (If a (i) Details of Investigato	e, (If any):ny):ry):	Version	number:
Acronym/ Short title (h) Protocol number (If a (i) Details of Investigato Name	e, (If any):ny):ry):	Version Department and	number:
Acronym/ Short title (h) Protocol number (If a (i) Details of Investigato Name	e, (If any):ny):ry):	Version Department and	number:
Acronym/ Short title (h) Protocol number (If a (i) Details of Investigato Name	e, (If any):ny):ry):	Version Department and	number:
Acronym/ Short title (h) Protocol number (If a (i) Details of Investigato Name	Designation and Qualification	Version Department and	number:
Acronym/ Short title (h) Protocol number (If a (i) Details of Investigator Name Principal Investigator/G	Designation and Qualification	Version Department and	number:
Acronym/ Short title (h) Protocol number (If a (i) Details of Investigator Name Principal Investigator/G	Designation and Qualification	Version Department and	number:
Acronym/ Short title (h) Protocol number (If a (i) Details of Investigator Name Principal Investigator/G	Designation and Qualification	Version Department and	number:
Acronym/ Short title (h) Protocol number (If a (i) Details of Investigator Name Principal Investigator/G	Designation and Qualification suide	Version Department and	number:

2.	FUN	IDING DETAILS	AND BUDGET				
	(a) 1	Total estimated	budget for site:				
		At site		In India	Globall	y	
	(b)	Self-funding \Box	l Institutio	nal funding 🔲 Fundin	g agency (Specif	; _у) 🗆	
	• •			•			
	•••••						
		SI	ECTION B	- RESEARCH RELA	TED INFO	RMATION	
3.		ERVIEW OF RES		ds):			
	• •						

	(b)	Type of study:					
		Basic Sciences		Clinical		Cross Sectional	
		Retrospective		Epidemiological/		Case Control	
		Prospective		Public Health		Cohort	
		Qualitative		Socio-behavioural		Systematic Review	
		Quantitative		Biological samples/ Data			
		Mixed Method		Any others (Specify)			
4.	MET	HODOLOGY					
	(a)	Sample size/ n	umber of partic	ipants <i>(as applicable)</i>			
		At site		In India	Globall	y	
		Control group.		Study	group		
		Justification fo	r the sample siz	e chosen (100 words); In case	e of qualitative s	tudy, mention the criteria	used for
		saturation					
³Su	mmari	ize in the simplest p	ossible way such tha	t a person with no prior knowledge of	the subject can easi	ly understand it.	

	(b)	Is the	ere an externa	l labor	atory/o	utsourcing invo	olved for investi	gations?	4 Yes □ No □	NA 🗆
(c) How was the scientific quality of the study assessed?										
		Indep	oendent exteri	nal rev	view 🛚	Review by sp	oonsor/Funder		Review within PI's institution	
			ew within mult arch group	i-cent	re 🗆	No review				
		Date	of review:						dd mm yy	
		Comr	ments of scien	ntific c	ommitte	ee, if any (100	words)			
									NFORMATION	
5.	REC	RUITI	MENT AND RE	SEAR	CH PAR	TICIPANTS				
	(a)	Туре	of participants	in the	e study:					
		Healt	thy volunteers	; 		Patients 🛚	Vulnerable p	ersons/	Special groups	
		Othe	ers		(Specify)					
		Who	will do the re	cruitm	nent?					
		Parti	cipant recruit	ment ı	methods	s used:				
Posters/									iends 🗆 Telephone 🗆	
		Othe	ers		(Specify))				
	(h)	i W	/ill there he vi	ılneral	hle ners	ons / special d	roups involved	?	Yes □ No □	1 να Π
	(5)					sons / special g		•	163 🗀 110 🗀	INAL
			hildren under			sons / special s	угоира	Pregna	nt or lactating women	П
			ifferently able			vsical)	П	_	yees/Students/Nurses/Staff	П
			lderly	(1.10		yordary			ionalized	
			conomically a	nd soc	cially dis	sadvantaged			es/Migrants/Homeless	
						rare diseases)			, ,	
		Α	ny other <i>(Spe</i>	cify):						
					or inclu	sion/exclusion				
		iv. A	re there any a	dditio	nal safe	guards to prote	ect research par	ticipants	?	

(c)	Is there any reimbursement to the participants?									
	If yes,	Monetary 🗆	Non-monetary 🛘	Provide	e details					
(d)		ere any incentives to	o the participants? Non-monetary	Provide	e details			Yes □ No □		
(e)		Monetary	ecruitment fees/ incenti Non-monetary				the PI / Ins	Yes No No		
6. BE	NEFITS	AND RISKS								
(a)		there any anticipate es, categorize the lev	d physical/social/psychorel of risk ⁵ :	ological disc	omforts,	/ risk to p	articipants	? Yes□ No□		
	Less	s than Minimal risk		Minima	ıl risk					
			imal risk or low risk ement strategy:				r high risk			
(b)	 What a	re the potential bene	efits from the study?	Yes	No	If yes,	Direct	Indirect		
	For the	e participant								
	For the	e society/community								
	For im	provement in science	e							
	Please		enefits justify the risks							
(c)		rerse events expecte					Yes	:		
			nd management strategi					Yes No		
		CONSENT								
(a)	Are you	ı seeking waiver of c	onsent? If yes, please sp	ecify reason	s and sk	ip to item	no. 8	Yes 🗆 No 🗆		
⁵For ca			ical Guidelines for Biomedical &							

(b)) Version number and date of Participant Information Sheet (PIS):										
	Version number an	d date of I	nformed Consent Forr	n (ICF):						
(c)	Type of consent pl	anned for :									
	Signed consent		Verbal/Oral consent		Witnessed co	onsent		Audio-Video (AV) consent			
	Consent from LAR (If so, specify from		For children<7 yrs parental/LAR consent		Verbal assent minor (7-12 y with parental	rs) along		Written assent from minor (13-18 yrs) alon with parental consent			
	Other										
(d)	Who will obtain the	informed	consent?								
	PI/Co-I \square Nurse/Counselor \square Research Staff \square Other \square (Specify)										
	Any tools to be use	ed									
(e)	Participant Informa	ation Sheet	(PIS) and Informed C	onsen	t Form (ICF)						
	English \square	Local lang	guage 🗆	Othe	r 🛘 (Specify).						
	List the languages	in which tr	anslations were done								
	If translation has no	ot been do	ne, please justify								
(f)	Provide details of o	consent red	quirements for previou	ısly sto	ored samples i	f used in t	he s	tudy ⁷			
	•••••										
(g)	Elements contained	in the Part	cicipant Information Sh	neet(P	IS) and Inform	ed Conse	nt F	orm (ICF)			
	Simple language		Data/ Sample sharing				-	related injury			
	Risks and discomforts Alternatives to participations	ation \square	Need to recontact Confidentiality			nt that con: cialization/		· —			
	Right to withdraw		Storage of samples					olves research			
	Benefits		Return of research resu	Its 🗆			•	ntifying data			
	Purpose and procedure Others(Specify)		Payment for participati	on 🗆		information y of EC	of P	I and Member			
8. PA	YMENT/COMPENSAT	ION									
(a)	Who will bear the co	osts related	to participation and	proce	dures ⁸ ?						
	PI 🗆	I	Institution	Sp	oonsor 🗆	Other a	agen	cies 🛘 (specify)			
(b)	Is there a provision	for free tre	eatment of research re	lated	injuries?			Yes □ No □ N/A	Α 🗆		
	If yes, then who wil	I provide t	he treatment?								
(c)	Is there a provision	Yes □ No □ N/	Ά 🗆								
	Sponsor 🗆 Ins	stitutional/	Corpus fund \Box	Projec	t grant 🛚	Insurar	ice				
(d)	Is there any provisio	n for medi	cal treatment or mana	geme	nt till the relat	edness is	dete	rmined for injury to th	ne		
	participants during	the study	period? If yes, specify					Yes □ No □ N/	Ά□		
(e)	Is there a provision	for ancilla	ry care for unrelated il	Iness	during the stu	dy period	? If y	es, please specify.	•••••		
								Yes 🗆 No 🗀 N/			
	ation on re-consent require age 54 in Section 5.8.	ments can be	found at National Ethical Gu	iidelines	s tor Biomedical ai	nd Health Re	searci	n Involving Human Participar	nts		

9.	. STORAGE AND CONFIDENTIALITY	
	(a) Identifying Information: Study Involves samples/data. If Yes, specify	Yes ☐ No ☐ NA ☐
	Anonymous/Unidentified \square Anonymized: Reversibly coded \square Irreversibly α	coded \square Identifiable \square
	If identifiers must be retained, what additional precautions will be taken to ensure	that access is limited /data is
	safeguarded? (e.g. data stored in a cabinet, password protected computer etc.)	
	(b) Who will be maintaining the data pertaining to the study?	
	(c) Where will the data be analyzed ⁹ and by whom?	
	(d) For how long will the data be stored?	
	(e) Do you propose to use stored samples/data in future studies?	Yes □ No □ Maybe □
	If yes, explain how you might use stored material/data in the future?	
	SECTION D: OTHER ISSUES	
	SECTION D. OTTER 1330ES	
10.	D. PUBLICATION, 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 participation	pants, if effective, once the
	study has finished? If yes describe in brief (Max 50 words)	Yes □ No □ NA □
	(d) Is there any plan for post research benefit sharing with participants? If yes, specify	Yes □ No □ NA □
	(e) Is there any commercial value or a plan to patent/IPR issues? If yes, please provide	e details Yes 🗆 No 🗆 NA 🗆
	(f) Do you have any additional information to add in support of the application, which	n is not included elsewhere in
	the form? If yes, provide details.	Yes □ No □

SECTION E: DECLARATION AND CHECKLIST 10

11. DI	ECLARATION (Please tick as applicable)										
	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/We confirm that this study will be conducted in accordance with the latest ICMR National Ethical Guidelines for Biomedical and Health Research Involving Human Participants and other applicable regulations and guidelines.										
	I/We confirm that this study will be conducted in accordance with the Drugs and Cosmetics Act 1940 and its Rules 1945 as amended from time to time, GCP guidelines and other applicable regulations and guidelines.										
	I/We will comply with all policies and guidelines of the institute and affiliated/collaborating institutions where this study will be conducted.										
	I/We will ensure that personnel performing this study are qualified, appropriately trained and will adhere to the provisions of the EC approved protocol.										
	I/We declare that the expenditure in case of injury related to the study will be taken care	of.									
	I/We confirm that an undertaking of what will be done with the leftover samples is provid	ded, if applicable.									
	I/We confirm that we shall submit any protocol amendments, adverse events report, significant deviations from protocols, progress reports and a final report and also participate in any audit of the study if needed.										
	I/We confirm that we will maintain accurate and complete records of all aspects of the st	udy.									
	I/We will protect the privacy of participants and assure confidentiality of data and biolog	gical samples.									
	I/We hereby declare that I/any of the investigators, researchers and/or close relative(s), have no conflict of interest (Financial/Non-Financial) with the sponsor(s) and outcome of study.										
	1/We have the following conflict of interest (PI/Co-I): 1										
	I/We declare/confirm that all necessary government approvals will be obtained as per receiver applicable.	quirements wherev-									
Na	me of PI:										
Sig	gnature:	dd mm yy									
Na	me of Co-PI:										
Sig	Signature: dd mm yy										
Na	Name of Guide:										
Sig	gnature:	dd mm yy									
Na	me of HOD:										
Sig	gnature:	dd mm yy									

¹⁰These 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)

12. CH	ECKLIST											
S. No	S. No Items Yes No NA Enclosure EC Remarks (If applicable)											
ADMI	NISTRATIVE REQUIREM	ENTS	5									
1	Cover letter											
2	Brief CV of all Investigato	rs										
3	Good Clinical Practice (GO	CP) tr	aining	of investi	gators in	last 3 years						
4	Approval of scientific com	nmitte	ee									
5	EC clearance of other cen	ters*										
6	Agreement between colla	borat	ing pa	rtners*								
7	MTA between collaboratin	ng pai	rtners*	•								
8	Insurance policy/certificat	:е										
9	Evidence of external labor outsourced laboratory stu					externally						
10	Copy of contract or agreem	ent sig	gned w	ith the spo	onsor or d	onor 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											
PROPOSAL RELATED												
12	Copy of the detailed prote	ocol ¹¹										
13	Investigators Brochure (If	appli	cable f	or drug/b	iological	s/device trials)						
14	Participant Information St Form (ICF)(English and tr			nd Partic	ipant Info	rmed Consent						
15	Assent form for minors (12	2-18 y	ears) ((English a	nd Transl	ated)						
16	Proforma/Questionnaire / Guides for Focused Group											
17	Advertisement/material to	o recr	uit par	rticipants	(fliers, po	osters etc)						
PERMI	SSION FROM GOVERNII	NG A	итно	RITIES								
	Other permissions	Requ	iired	Not required	Received	Applied dd/ mm/yy				EC Remarks		
18	CTRI]									
19	DCGI		1									
20	HMSC		1									
21	NAC-SCRT											
22	ICSCR		1									
23	RCGM]									
24	4 GEAC 🗆 🗆											
25												
26												
27	Others (Specify)											
ANY O	THER RELEVANT INFOR	RMAT	TON/	DOCUME	ENTS RE	LATED TO TH	IE STU	DY				
	Item		YES	NO	NA	Enclosure no.				EC remarks		
28												
29	29											

*For multicentre 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

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

Version 2.0

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