

## **Application Form for Initial Review**

Institute Ethics Committee Indian Institute of Technology, Delhi *Ph: 011-26591221* 

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**

(a) Name of Organization	າ:		
(b) Name of Ethics Con	nmittee:		
(c) Name of Principal Inv	estigator:		
(d) Department/Division:		(e) Date of submis	sion: dd mm yy
(f) Type of review requeste	ed¹:		
Exemption from review	Expedited rev	iew 🗆 Full com	mittee review
g) Title of the study:			
4 (01	(If any):		
Acronym/ Short title,	(II ally)		
-			number:
h) Protocol number (If a	any):		
h) Protocol number (If a	any):		
h) Protocol number (If a	Designation and Qualification	Version  Department and	number:
h) Protocol number (If a i) Details of Investigators: Name	Designation and Qualification	Version  Department and	number:
h) Protocol number (If a i) Details of Investigators: Name	Designation and Qualification	Version  Department and	number:
h) Protocol number (If a i) Details of Investigators: Name	Designation and Qualification	Version  Department and	number:
h) Protocol number (If a i) Details of Investigators: Name	Designation and Qualification	Version  Department and	number:
h) Protocol number (If a i) Details of Investigators: Name Principal Investigator/Gu	Designation and Qualification	Version  Department and	number:
(h) Protocol number (If a (i) Details of Investigators: Name Principal Investigator/Gu	Designation and Qualification	Version  Department and	number:
h) Protocol number (If a i) Details of Investigators: Name Principal Investigator/Gu	Designation and Qualification	Version  Department and	number:
h) Protocol number (If a i) Details of Investigators: Name Principal Investigator/Gu	Designation and Qualification ide	Version  Department and	number:

2.	FUN	IDING DETAILS	AND BUDGET											
	(a)	Total estimate	d budget for si	te:										
		At site		In India	Globally									
	(b)	Self-funding $\square$	cify) 🗆											
		SI	ECTION B	- RESEARCH F	RELATED INFO	ORMATION								
3.	OVERVIEW OF RESEARCH  (a) Lay summary <sup>3</sup> (within 300 words):													
							•••••							
							•••••							
	(b) <sup>-</sup>	Гуре of study:												
		Basic Sciences		Clinical		Cross Sectional								
				Epidemiological/	_	Case Control	_							
		Prospective		Public Health		Cohort								
		Qualitative		Socio-behavioural		Systematic Review								
		Quantitative		Biological samples/ D	ata 🗆									
		Mixed Method		Any others (Specify)										
4.	MET	HODOLOGY												
	(a)	Sample size/ no	umber of particip	oants (as applicable)										
					Globally									
					•									
						e study, mention the criteria								
		saturation												
³Sι	ımmar	rize in the simplest po	ossible way such tha	t a person with no prior knowl	edge of the subject can ea	sily understand it.								

(b)	Is there an external laboratory/outsourcing inv	volved for investi	gations? <mark>4</mark>	Yes ☐ No l	$\square$ NA $\square$
(c)	How was the scientific quality of the study ass	essed?			
	Independent external review $\square$ Review by	sponsor/Funder		Review within PI's institution	n 🗖
	Review within multi-centre    No review research group				
	Date of review:			dd mm yy	
	Comments of scientific committee, if any (100	) words)			
	SECTION C: PARTICIP	PANT RELA	TED IN	IFORMATION	
i. RE	CRUITMENT AND RESEARCH PARTICIPANTS				
(a)	Type of participants in the study:				
	Healthy volunteers ☐ Patients ☐	Vulnerable p	ersons/ S	pecial groups $\Box$	
	Others				
	Who will do the recruitment?				
	Participant recruitment methods used:				
	Posters/	Patients / Fa		nds  Telephone	
	Others				
(b)	i. Will there be vulnerable persons / special	groups involved	?	Yes □ No	□ NA □
	ii. If yes, type of vulnerable persons / special	groups			
	Children under 18 yrs		Pregnar	t or lactating women	
	Differently abled (Mental/Physical)		Employe	ees/Students/Nurses/Staff	
	Elderly		Institutio	nalized	
	Economically and socially disadvantaged		Refugee	s/Migrants/Homeless	
	Terminally ill (stigmatized or rare diseases)				
	Any other (Specify):				
	iii. Provide justification for inclusion/exclusion	usion			
					iv.
	Are there any additional safeguards to protect re	esearch participan	its?		

	(c)	Is ther	Yes 🗆	No 🗖								
		If yes,	Monetary C		Non-monetary		Provide	details				
	(d)		ere any incen	_	participants?		Provide	details			Yes 🗆	No 🗆
	(0)		oro ony porti	oinant roor	uitmont food/inc		for the of		wided to	the DI / Inet	itution?	
	(e)		Monetary C	_	uitment fees/ ind		Provide			e P1/ IIISu	Yes 🗆	No 🗆
6.	BEN	NEFITS A	AND RISKS									
	(a)		there any an		hysical/social/ps f risk <sup>5</sup> :	sycholog	ical disco	omforts/	risk to p	articipants?	Yes 🗖	No 🗆
		Less	s than Minima	ıl risk			Minimal	risk				
		Mino	or increase ov	er minimal	risk or low risk		More th	an minii	mal risk c	r high risk		
		ii. Des	cribe the risl	k manager	nent strategy:							
	(b)	 What aı	e the potenti	al benefits	from the study?		Yes	No	If yes,	Direct	Indirect	
		For the	participant									
		For the	society/com	nmunity								
		For imp	provement in	science								
		Please	describe h	now the b	enefits justify	the risk	(S					
	(c)	Are adv	erse events e	expected in	the study <sup>6</sup> ?					Yes	□ No □ N	IA 🗆
		Are rep	orting proced	ures and m	anagement stra	tegies d	escribed i	in the s	tudy?		Yes $\square$	No 🗆
		If Y	es, Spe	cify								
7.	INF	ORMED	CONSENT									
	(a)	Are you	seeking waiv	er of cons	ent? If yes, pleas	se speci	fy reason:	s and sk	kip to iten	n no. 8	Yes 🗆	No 🗖
<sup>5</sup> F	or cat	egories of			uidelines for Biomedi							

	(b)	b) Version number and date of Participant Information Sheet (PIS):								
		Version number and	d date of	Informed Consent Fo	rm (l	CF):				
	(c)	Type of consent plan	nned for :							
		Signed consent		Verbal/Oral consent		Witnessed co	onsent	_	Audio-Video (AV)	
		Consent from LAR (If so, specify from v	□ vhom)	For children<7 yrs parental/LAR consent		Verbal asser minor (7-12 y with parental	yrs) along		Written assent from Iminor (13-18 yrs) along with parental consent	
		Other								
		(specify)								
	(d)	Who will obtain the in	nformed (	consent?						
	` ,	PI/Co-I ☐ Nur	se/Couns	elor  Research						
		-								
	(e)	· <u>_</u>		: (PIS) and Informed C juage □		, ,	1			
		•	•	,		` ' ' ' '				
	(f)	Provide details of co	nsent red	quirements for previous	sly sto	ored samples	if used in t	he st	udy <sup>7</sup>	
		Elements contained in Simple language Risks and discomforts Alternatives to participat Right to withdraw Benefits Purpose and procedure Others(Specify)		ticipant Information Sh Data/ Sample sharing Need to recontact Confidentiality Storage of samples Return of research res Payment for participatio	C Ults	Comper Stateme Comme Stateme Use of Contact	nsation for ent that cons ercialization/ ent that stud photograph	study ent is Bene ly inv	related injury  voluntary	
8.	PA	MENT/COMPENSAT	ION							
	(a)	Who will bear the cos	ts related	to participation and p	rocec	ures <sup>8</sup> ?			_	
		PI 🗆	I	Institution	Sı	oonsor $\square$	Other a	geno	cies 🛘 (specify)	
	(b) Is there a provision for free treatment of research related injuries?  Yes □ No □ N/A □  If yes, then who will provide the treatment?									
	(c)	Is there a provision for	or compe	nsation of research rel	ated S	SAE? If ye	es, specify.		Yes ☐ No ☐ N/A ☐	
Sponsor ☐ Institutional/Corpus fund ☐ Project grant ☐ Insurance ☐										
	(d)	Is there any provision	for medi	cal treatment or mana	geme	nt till the relat	edness is	dete	rmined for injury to the	
				period? If yes, specify.					Yes No No N/A	
	(e)			ry care for unrelated ill						
									Yes ☐ No ☐ N/A ☐	
201	7, Pa	ge 54 in Section 5.8.			idelines	for Biomedical ar	nd Health Res	earch	Involving Human Participants	
<u>_[]</u>	liose	undertaking from PI confirm	miy ule san	IIC .					Version 2.0 05	

9.	(a) Identifying Information: Study Involves samples/data. <i>If Yes, specify</i> Anonymous/Unidentified □ Anonymized: Reversibly coded □ Irreversibly coded identifiers must be retained, what additional precautions will be taken to ensure that as safeguarded? (e.g. data stored in a cabinet, password protected computer etc.)	ccess is limited /data is
	saleguarded: (e.g. data stored in a cabinet, password protected computer etc.)	
	Who will be maintaining the data pertaining to the study?	(c)
	(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?	/es ☐ No ☐ Maybe ☐
	SECTION D: OTHER ISSUES	
10.	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 participants, i study has finished? If yes describe in brief (Max 50 words)	f effective, once the  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 details	Yes No NA
	(f) Do you have any additional information to add in support of the application, which is not the form? If yes, provide details.	included elsewhere in Yes ☐ No ☐

## SECTION E: DECLARATION AND CHECKLIST 10

11. C	DECLARATION (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 provided, 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 study.										
	I/We will protect the privacy of participants and assure confidentiality of data and biological 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										
<u> </u>	I/Ma dealers/confirm that all passagery government approvals will be obtained as per requirements wherever										
	I/We declare/confirm that all necessary government approvals will be obtained as per requirements wherever applicable.										
Na	ime of PI:										
Sig	gnature:dd mm yy										
Na	Name of Co-PI:										
Siç	Signature: dd mm yy										
Na	Name of Guide:										
Siç	Signature: dd mm yy										
Na	me of HOD:										
Sig	gnature: dd mm yy										

<sup>&</sup>lt;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)

12. CH	12. CHECKLIST										
S. No	S. No Items								NA	Enclosure No	EC Remarks (If applicable)
ADMI	NISTRATIVE REQUIREM	IENT	S							140	(п аррисало)
1	Cover letter										
2	Brief CV of all Investigato	rs									
3	Good Clinical Practice (G	g of investi	last 3 years								
4	Approval of scientific con	nmitte	ее								
5	EC clearance of other cen	iters*									
6	Agreement between colla	abora	ting pa	artners*							
7	MTA between collaboratii	ng pa	rtners	*							
8	Insurance policy/certification	te									
9	Evidence of external labo outsourced laboratory st					n externally					
10	Copy of contract or agreem	ent si	gned v	vith 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										
PROP	OSAL RELATED										
12	Copy of the detailed proto	ocol <sup>11</sup>									
13	Investigators Brochure (If	appl	icable	for drug/b	oiological	s/device trials)					
14	Participant Information SI Form (ICF)(English and to			and Partic	ipant Info	ormed Consent					
15	Assent form for minors (12	2-18 չ	/ears)	(English a	nd Trans	lated)					
16	Proforma/Questionnaire / Guides for Focused Group										
17	Advertisement/material to	o recr	uit pa	rticipants	(fliers, po	osters etc)					
PERMI	SSION FROM GOVERNI	NG A	UTHO	DRITIES							
	Other permissions	Req	uired	Not required	Receive	d Applied dd/ mm/yy				EC Remarks	
18	CTRI	[	o j				ĺ				
19	DCGI	[	o i								
20	HMSC	[	_								
21	NAC-SCRT	[	_								
22	ICSCR	[									
23	RCGM	[									
24	GEAC										
25	BARC										
26	6 Tribal Board										
27	Others (Specify)										
ANY O	THER RELEVANT INFO	RMA	TION/I	DOCUME	NTS RE	LATED TO TH	E STU	DY			
	Item		YES	NO	NA	Enclosure no.				EC remarks	-
28											
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

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

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