

## Application Form for Initial Review

## **Central University of South Bihar**

SH-7, Gaya - Panchanpur Road, Village - Karhara, Post-Fatehpur P.S. - Tekari, District - Gaya (Bihar) Pin- 824236

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	_S		
(a) Name of Organization:			
(b) Name of Ethics Committee	:		
(c) Name of Principal Investig	jator:		
(d) Department/Division:		(e) Date of submi	ission: dd mm yy
f) Type of review request	red¹:		
Exemption from revie			nmittee review 🗆
Actoriyin/ Short duc, (ii	αιιγ)		
(h) Protocol number (If any):		Version n	umher:
		Version n	umber:
(i) Details of Investigator	s:		
		Department and Institution	Address for communication <sup>2</sup>
(i) Details of Investigator	s:  Designation and Qualification	Department and	
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(i) Details of Investigator	s:  Designation and Qualification	Department and	
i) Details of Investigator Name Principal Investigator/G	Designation and Qualification	Department and	
i) Details of Investigator Name	Designation and Qualification	Department and	
Name Principal Investigator/G	Designation and Qualification	Department and	
Name Principal Investigator/G	Designation and Qualification	Department and	
Name Principal Investigator/G	Designation and Qualification	Department and	
Name Principal Investigator/G  Co-investigator/student	Designation and Qualification uide	Department and	
Name Principal Investigator/G  Co-investigator/student  (j) Number of studies who	Designation and Qualification uide	Department and Institution	

Refer to National Ethical Guidelines for Biomedical and Health Research Involving Human Participants 2017 on Page 36 Table 4.2. for types of review Version 2.0

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

2.	FUN	DING DETAILS	AND BUDGET									
	(a)	Total estimated bu	dget for site:									
				In India								
	(b) 5	b) Self-funding $\square$ Institutional funding $\square$ Funding agency (Specify) $\square$										
		SI	ECTION B	- RESEARCH RELA	TED INFORI	MATION						
3	OVE	ERVIEW OF RES	FARCH									
٠.												
							•••••					
	(h) 7											
		Type of study:			П		_					
		Basic Sciences	_	Clinical		Cross Sectional  Case Control						
		Retrospective Prospective		Epidemiological/ Public Health	Ц	Cohort	П					
		Qualitative		Socio-behavioural		Systematic Review						
		Quantitative		Biological samples/ Data		5,500	_					
		Mixed Method		Any others (Specify)								
4.	MET	HODOLOGY										
	(a)	Sample size/ i	number of part	icipants (as applicable)								
		At site		In India	Globally							
		Control group		Study	group							
		Justification for	the sample size	chosen (100 words); In case of	of qualitative study	mention the criteria use	ed for					
		saturation										
							•••••					
³Sı.	ımmar	ize in the simplest p	ossible wav such tha	rt a person with no prior knowledge o	f the subject can easily	understand it.	•••••					
		, , , , , , , , , , , , , , , , , , ,	,	,								

(b)	Is there an external laborat	4 Yes □ No □	Yes □ No □ NA □							
(c)	) How was the scientific quality of the study assessed?									
	Independent external revie	w 🛘 Review by sp	onsor/Funder		Review within PI's institution					
	Review within multi-centre research group	☐ No review								
	Date of review:				dd mm yy					
	Comments of scientific cor	nmittee, if any (100	words)							
	SECTION	C: PARTICIPA	ANT RELA	TED	NFORMATION					
REC	CRUITMENT AND RESEARCH	H PARTICIPANTS								
(a)	Type of participants in the s	tudy:								
	Healthy volunteers $\Box$	Patients $\square$	Vulnerable p	ersons/	Special groups $\square$					
	Others	cify)								
	Who will do the recruitment?									
	Participant recruitment me	ethods used:								
	leaflets/Letters Sc	V/Radio ads/ Docial media/stitution website	Patients / Fa		ends   Telephone					
	Others	ecify)								
(b)	i. Will there be vulnerable	persons / special g	roups involved	?	Yes □ No □	l na □				
	ii. If yes, type of vulnerable	e persons / special (	groups							
	Children under 18 yrs			Pregna	nt or lactating women					
	Differently abled (Ment	al/Physical)		Employ	/ees/Students/Nurses/Staff					
	Elderly			Institu	tionalized					
	Economically and socia	lly disadvantaged		Refuge	es/Migrants/Homeless					
	Terminally ill (stigmatiz	ed or rare diseases)								
	Any other (Specify):		<b></b>							
		clusion/exclusion								
		,								
	iv. Are there any additional	safeguards to protect	research particip	ants?						

(c	c) Is there any reimbursemen	Yes 🗆 No 🗆					
	If yes, Monetary $\square$	Non-monetary 🗖	Provide	details			
(c	d) Are there any incentives to	o the participants?	Provide	details			Yes □ No □
(e	e) Are there any participant r	ecruitment fees/ incentiv	es for the s	study pro	ovided to	the PI / Ins	titution?
	If yes, Monetary 🗆	Non-monetary 🗆	Provide	details			Yes 🗌 No 🗖
6. BI	ENEFITS AND RISKS						
(a	i. Are there any anticipate		ogical disc	omforts	/ risk to p	articipants	? Yes □ No □
	Less than Minimal risk		Minima	ıl risk			
	Minor increase over mini ii. Describe the risk managem					r high risk	
(b	o) What are the potential bene	fits from the study?	Yes	No	If yes,	Direct	Indirect
	For the participant						
	For the society/community	,					
	For improvement in scienc	e					
	Please describe how the benefits	justify the risks					
(c	:) Are adverse events expected						□ No □ NA □
``	Are reporting procedures and If Yes, Specify	nd management strategies					Yes  No
	NFORMED CONSENT						
(a	a) Are you seeking waiver of c	onsent? If yes, please spe	cify reason	s and sk	ip to item	no. 8	Yes 🗆 No 🗖
5For o	categories of risk refer to National Eth	ical Guidelines for Biomedical &					

(b)	Version number and d	ate of Par	ticipant Information Shee	et (PIS):				
	Version number and da	ate of Info	rmed Consent Form (ICF):					
(c)	Type of consent pla	nned for	:					
	Signed consent		Verbal/Oral consent		Witnessed consen	t 🗆	Audio-Video (AV) consent	
	Consent from LAR (If so, specify from v		For children<7 yrs parental/LAR consent		Verbal assent from minor (7-12 yrs) a with parental cons	long	Written assent from minor (13-18 yrs) alo with parental consent	
	Other							
(d)	Who will obtain the i	nformed	consent?					
	-	-						
(e)	,		t (PIS) and Informed C					
(E)	· <u> </u>		guage $\square$					
	_							
(f)	Provide details of co	nsent re	quirements for previo	usly st	ored samples if use	d in the	study <sup>7</sup>	•••••
(g)	Elements contained i Simple language Risks and discomforts Alternatives to participa Right to withdraw Benefits Purpose and procedure Others(Specify)		ticipant Information S Data/ Sample sharing Need to recontact Confidentiality Storage of samples Return of research re Payment for participat	C C Sults	Compensation Statement th Commercializ Statement tha Use of photo	n for stu at conse ation/ E at study i graphs/ mation o	dy related injury  ent is voluntary  Benefit sharing  involves research  Identifying data  If PI and Member	
8. <b>PA</b> `	YMENT/COMPENSATION	ON						
			ed to participation an	d prod	edures <sup>8</sup> ?			
	PI 🗆		Institution $\square$	Sı	oonsor 🛭 Ot	herage:	encies 🛭 (specify)	
(b)	·		eatment of research re		-		Yes □ No □ N/	а <b>П</b>
(c)			ensation of researchr /Corpus fund			ecify. surance	Yes □ No □ N/#	A 🗆
(d)			dical treatment or mar period? If yes, specify	_	ent till the relatedn	ess is d	etermined for injury to Yes □ No □ N/	
(e)	Is there a provision f	or ancilla	ary care for unrelated i	llness	during the study pe	eriod? If	yes, please specify.	а 🗆

9.	STORAGE AND CONFIDENTIALITY		
	(a) Identifying Information: Study Involves samples/data. <i>If Yes, specify</i> Anonymous/Unidentified □ Anonymized: Reversibly coded □ Irreversibly coded □  If identifiers must be retained, what additional precautions will be taken to ensure that accomplying the product of the	cess is limited /data is	] s
	safeguarded? (e.g. data stored in a cabinet, password protected computer etc.)		
	(b) Who will be maintaining the data pertaining to the study?		
	(d) For how long will the data be stored?		
	-	s 🗆 No 🗆 Maybe 🗖	
	If yes, explain how you might use stored material/data in the future?		
	SECTION D: OTHER ISSUES		
10.	PUBLICATION, BENEFIT SHARING AND IPRISSUES		
	(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)	Yes □ No □ NA □	]
	(d) Is there any plan for post research benefit sharing with participants? If yes, <i>specify</i>	Yes 🗆 No 🗖 NA 🗀	]
	(e) Is there any commercial value or a plan to patent/IPR issues? If yes, please provide details	s Yes 🗆 No 🗆 NA 🗖	
	(f) Do you have any additional information to add in support of the application, which is not in the form? If yes, provide details.	ncluded elsewhere in Yes 🗖 No 🗖	

## SECTION E: DECLARATION AND CHECKLIST 10

11. 0	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 docu	ıments.									
	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 the provisions of the EC approved protocol.	l adhere to									
	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 ap	plicable.									
	I/We confirm that we shall submit any protocol amendments, adverse events report, significant d from protocols, progress reports and a final report and also participate in any audit of the study										
	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 samp	ples.									
	I/We hereby declare that I/any of the investigators, researchers and/or close relative(s), have no interest (Financial/Non-Financial) with the sponsor(s) and outcome of study.	conflict of									
	I/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 requirement er applicable.	nts wherev-									
Na	ame of PI:										
Sig	gnature:dd m	m yy									
Na	Name of Co-PI:										
Sig	Signature: dd mm yy										
Na	ame of Guide:										
Sig	gnature:dd m	m yy									
Na	ame of HOD:										
Sig	gnature:dd m	ım yy									

<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. CHECKLIST											
S. No		Item		Yes	No	NA	Enclosure No	EC Remarks (If applicable)			
ADMI	NISTRATIVE REQUIREM	IENT	S						<u> </u>	.,,0	( appeas.e)
1	Cover letter										
2	Brief CV of all Investigato	rs									
3	Good Clinical Practice (GC	CP) tr	aining	of investi	gators in	last 3 years					
4	Approval of scientific con	nmitt	ee								
5	EC clearance of other cen	ters*									
6	Agreement between colla	.bora	ting pa	ırtners*							
7	MTA between collaboratii	ng pa	rtners	*							
8	Insurance policy/certifica	te									
9	Evidence of external labo outsourced laboratory stu	rator udy Q	y credo A/QC	entials in c certificati	case of a on	n externally					
10	Copy of contract or agreem	ent s	igned v	with the sp	onsor or	donor agency					
11	Provide all significant p negative decision or m authorities for proposed s and modification(s) to pro	odifie study	ed pro (whetl	tocol) by	other	ECs/Regulatory					
PROPO	DSAL RELATED										
12	Copy of the detailed prot	ocol¹	1								
13	Investigators Brochure (If	appli	cable f	or drug/b	iological	s/device trials)					
14	Participant Information SI Form (ICF)(English and tra			nd Particip	ant Infor	med Consent					
15	Assent form for minors (	12-18	years	) (English	and Tran	islated)					
16	Proforma/Questionnaire , Guides for Focused Group										
17	Advertisement/material t	o rec	ruit pa	rticipants	(fliers, p	osters etc)					
PERMI	SSION FROM GOVERNI	NG A	UTHC	DRITIES							
	Other permissions	Req	uired	Not required	Receive	d Applied dd/ mm/yy				EC Remarks	
18	CTRI	[	<b>-</b>								
19	DCGI	[									
20	HMSC	[									
21	NAC-SCRT	[									
22	ICSCR	[									
23	RCGM	[	_								
24	GEAC 🔲 🗎										
25	BARC 🗆 🗆										
26	Tribal Board										
27	27 Others (Specify)										
ANY C	THER RELEVANT INFO	RMA	TION	/DOCUM	ENTS RI	ELATED TO TI	HE ST	UDY			
	ltem		YES	NO	NA	Enclosure no.				EC remarks	
28											
29	9										

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 08

<sup>\*</sup>For multicentre research.