

Institutional Ethics Committee Guru Gobind Singh Indraprastha University Sector 16-C, Dwarka, Delhi-110078

Dated : 07.07.2021

Notice

A meeting of the Institutional Ethics Committee will be held (online) on 19th July,2021 at 11:30 a.m. All the faculty members who wish to submit their research proposals for ethical clearance are requested to fill the prescribed format attached herewith. You are requested to submit the soft as well as hard copy of the filled format along with the proposal and cover letter to the USBT office latest by 15th July, 2021. Kindly attach all the mandatory documents as per the format. The checklist for the same is given in the point No. 12 of the format. The complete ICMR National Ethical Guidelines are available at the ICMR website.

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Dr. Rinu Sharma Member Secretary, IEC GGSIPU

Prof. K.K. Aggarwal Dean USBT

Copy to: Deans, all USS Head, UITS. Kindly upload the notice on University website

Despatch No. GGSIPU/USBT/ 8946

ICMR-Central Ethics Committee on Human Research (CECHR)

ICCM2 INDIAN COUNCIL OF MEDICAL RESEARCH	NCDIR NATIONAL CENTRE FOR DISEASE INFORMATICS AND RESEARCH
NCD Public Hea Collaborate Inn	th Actions and Policies

(Annexure 13)

Format for Curriculum Vitae for Investigators

EC Ref. No.(for office use):

Name:				
Present affiliation(Job title, departme	ent, and organisation):			
Address(Full work address):				
Telephone number:	Email address:			
Qualifications:				
Professional registration (Name of bo	ody, registration number and date of registration):			
Previous and other affiliations(Includ	le previous affiliations in the last 5 years and other current affiliations):			
Projects undertaken in the last 5 year	rs:			

Relevant research tr	raining/experience in the area ²⁵ :	
Relevant publication	ns (Give references to all relevant p	publications in the last five years):
		Date: Click here to enter a date
Signature		Date: Click here to enter a date.
Signature		Date: Click here to enter a date.
Signature		Date: Click here to enter a date.
Signature		Date: Click here to enter a date.
Signature		Date: Click here to enter a date.
Signature		Date: Click here to enter a date.

²⁵Details of any relevant training in the design or conduct of research, for example in the Ethics Training, Human participants' protection courses, Clinical Trials Regulations, Good Clinical Practice, consent, research ethics training or other training appropriate to non-clinical research. Give the date of the training

Application Form for Initial Review						
Logo of the Institut	te	(Name of the li	nstitution)	EC Ref. No.(for office use):		
General Instructio	ns: a) Tick one or me b) Attach additio	ore as applicable. N mal sheets if requir		ot applicable		
	SEC	TION A - BASIC	INFORMA	TION		
c) Name of Prin	anization: Ethics Committee: ncipal Investigator:					
d) Department/			(e) Dat	e of Submission: Click here to enter a date.		
f) Type of revie Exemption fr	w requested ¹ : om Review	Expedited Revie	w 🗖	Full Committee Review 🗖		
g) Title of the st	tudy:					
Acronym/ Sh	ort title, (If any):					
h) Protocol num i) Details of Inv			Version r	umber:		
Name	Designation and Qualification	Department and Institution	Address fo	r communication ²		
Principal Investiga	ator/Guide					
Co-investigator/s	tudent/fellow					
j) Number of st	tudies where applicar	nt is a:				
i) Princip	oal Investigator at tim	e of submission:	ii) Co-Ir	vestigator at time of submission:		
k) Duration of t	he study:					
¹ Refer to National E	Ethical Guidelines for Biomed	ical & Health Research Invo	olving Human Pai	ticipants 2017on Page 36 Table 4.2. for the types of		
review ² Include telephone/I	mobile, fax numbers and emo	ail id				

a) Tot Ats		a G stitutional funding 🗖	lobally Fundir	ng agency	
			(Specify)		
	SECTION B	- RESEARCH RELAT	ED INFORI	MATION	
3. OV	ERVIEW OF RESEARCH				
(a)	Lay Summary of study ³ (within	300 words)			
(b)	Type of study: Basic Sciences RetrospectiveImage: Construct of the sector of the	Clinical Epidemiological/ Public Health Socio-behavioural Biological samples/Data Any others (Specify)		Cross Sectional Case Control Cohort Systematic Review	
4. ME (a)	THODOLOGY Sample size/ No. of Participant At site In India Control group Study Group Justification for the sample size used for saturation	a Globally	n case of qual	itative study, mentio	on the criteria
(b) (c)	Is there an external laboratory How was the scientific quality Independent external review Review within multi- centre research group		or investigati	ons? ⁴ Yes 🔲 No Review within Pl's institution	
	Date of review: Comments of Scientific Comm	ittee, if any(100 words)		Click here to	enter a date.
	narize in the simplest possible way such that ticipant samples are sent outside for investi				

etc.

		65.67							
REC	CRUITME		ION C - PARTICIPAN	-	KELA I				
(a)		participants				nerable person/ cial groups		Others (Specify)	
	-	ill do the recr ant recruitme	uitment? ent methods used:						
	Poster leaflet	rs/ [ss/Letters	TV/Radio ads/Social media/Institu website	tion		Patients / Family/Friends visiting hospitals		Telephone	
	Others	s(Specify)							
(b)	i. ii.		vulnerable persor f vulnerable persor		•	nvolved? Ye	es 🗖	No 🗖 NA 🛛	
		Children und	er 18 yrs			Pregnant or lact	ating v	vomen	
		Differently at	oled (Mental/Physi	cal)		Employees/Stud	ents/N	Nurses/	
		Elderly				Staff Institutionalized			
		-	and socially disady (stigmatized or rar	-		Refugees/Migra	nts/Hc	omeless	
		Any other (Sp	ecify):						
	iii.	Provide justi	fication for inclusic	on/exclusio	n				
	iv.	Are there an	y additional safegu	ards to pro	otect re	esearch participant	:s?		
(c)			sement to the part					Yes 🗖 🛚	No 🗖
	If yes,	Monetary	Non-monetary	Pro ⁻	vide de	etails			
(d)	Are the	ro any incont	ives to the particip	ant?				Yes 🗖	
(-)				_					
	If yes,	Monetary	Non-monetary	📙 Provi	de det	ails			
:)	Are the	re any partici	pant recruitment f	ees/ incent	ives fo	r the study provid	ed to t	he PI/ Institut	ion?
	lf yes,	Monetary	Non-monetary	D Provid	le deta	ils		Yes 🗖 No	
		·	,						

									Yes 🔲 No	o 🗖	
	If yes, categorize Less than Minir				Minima	al risk					
	Minor increase Low Risk ii. Describe the risk				∕lore tl	han Minim	al Ris	k or H	igh Risk		
(b)	What are the potent	ial be	nefits from the st	udy?	Yes	No If y	'es,	Direc	t In	direct	
	For the participant								1		
	For the society/com	munit	Y						1		
	For improvement in	scien	ce						1		
	Please describe how			risks					4		
(c)	Are Adverse Events e	exnec	ted in the study ⁶ ?						Yes 🗖 No		
		•	,								
	Are reporting proced If Yes, Specify	dures	and management	strate	gies de	escribed in	the s	tudy?	Yes 🗖 N	o 🗖	
7. 11		dures	and management	strate	gies de	escribed in	the s	tudy?	Yes 🗖 N	ο 🗖	
7. II (a)	If Yes, Specify		-		-						
	If Yes, Specify NFORMED CONSENT Are you seeking waiw Version number and	ver of date	consent? If yes, p of Participant Info	lease s	specify on She	reasons ar et (PIS):					
(a)	If Yes, Specify NFORMED CONSENT Are you seeking waiv	ver of date date	consent? If yes, p of Participant Info of Informed Cons	lease s	specify on She	reasons ar et (PIS):					
(a) (b)	If Yes, Specify NFORMED CONSENT Are you seeking waiv Version number and Version number and	ver of date date	consent? If yes, p of Participant Info of Informed Cons for : Verbal/ oral	lease s	specify on She rm (ICF	reasons ar et (PIS): ⁻): 'itnessed			uestion 8. Yes Audio-Video	No	
(a) (b)	If Yes, Specify NFORMED CONSENT Are you seeking waiv Version number and Version number and Type of consent plan	ver of date date	consent? If yes, p of Participant Info of Informed Cons	lease s ormatio ent Fo	specify on She rm (ICF W co	reasons ar et (PIS): -):	nd ski	p to q	uestion 8. Yes	No	—]
(a) (b)	If Yes, Specify NFORMED CONSENT Are you seeking wain Version number and Version number and Type of consent plan Signed consent	ver of date date	of Participant Info of Informed Cons for : Verbal/ oral consent	lease s ormatio ent Fo	specify on She rm (ICF W co Ve frc 12 wi	reasons ar et (PIS): ⁻): 'itnessed nsent	nd ski t 7-	p to q	uestion 8. Yes Audio-Video (A/V) consent	No No t nt 13- with]
(a) (b) (c)	If Yes, Specify NFORMED CONSENT Are you seeking waiv Version number and Version number and Type of consent plan Signed consent Consent from LAR (If so, specify from whom) Other (specify)	ver of date date ined f	f consent? If yes, p of Participant Info of Informed Cons for : Verbal/ oral consent For children<7 yes parental/LAR consent	lease s ormatio ent Fo	specify on She rm (ICF W co Ve frc 12 wi	reasons ar et (PIS): -): /itnessed nsent rbal assent om minor (` yrs) along th parenta	nd ski t 7-	p to q	Audio-Video (A/V) consent Written Asser from Minor (1 18 yrs) along	No No t nt 13- with	
(a) (b)	If Yes, Specify NFORMED CONSENT Are you seeking waiw Version number and Version number and Type of consent plan Signed consent Consent from LAR (If so, specify from whom)	ver of date date ined f	f consent? If yes, p of Participant Info of Informed Cons for : Verbal/ oral consent For children<7 yes parental/LAR consent	lease s ormatio ent Fo	specify on She rm (ICF W co Ve frc 12 wi	reasons ar et (PIS): -): /itnessed nsent rbal assent om minor (` yrs) along th parenta	nd ski t 7-	p to q	Audio-Video (A/V) consent Written Asser from Minor (1 18 yrs) along	No No t nt 13- with	

(e)	English 🗖	Loca	neet(PIS) and Informe I language h translations were de		ent Form (ICF) other (<i>specify</i>)
(f)			n done, please justify t requirement for pre	viously	stored samples if used in the study ⁷
(g)	Elements contained	d in the	e Participant Informat	ion Sh	eet(PIS) and Informed Consent Form (ICF)
	Simple language		Data/ Sample sharing		Compensation for study related injury
	Risks and		Need to recontact		Statement that consent is voluntary
	discomforts Alternatives to participation		Confidentiality		Commercialization/benefit sharing
	Right to withdraw		Storage of samples		Statement that study involves research
	Benefits		return of research		Use of photographs/ identifying data
	Purpose and procedure		results Payment for participation		Contact information of PI and Member
	Others(Specify)				
8. P (a	AYMENT/COMPENS) Who will bear the PI	costs	related to participatic		procedures ⁸ ? onsor Dther agencies(specify)
(b) Is there a provisio	on for f	ree treatment of rese	arch re	elated injuries? Yes 🗖 No 🗖 NA 🗖
(c	•	•	vide the treatment? compensation of resea	arch re	lated SAE? If yes, specify. Yes 🗖 No 🗖 NA
	Sponsor 🗖 In	stitutic	n/ Corpus funds 🔲	P	Project grants 🔲 Insurance 🗖
(d			r medical treatment o s during the study per		agement till the relatedness is determined for yes, specify. Yes No NA
(e)	Is there a provision specify.	for and	illary care for unrelat	ed illn	ess during the study period? If yes, please Yes 🔲 No 🔲 NA
					Version 2.0 05

P	ation on re-consent requirements can be found at National Ethical Guidelines for Biomedical & Health Research Involving Human articipants 2017,Page 54 in Section 5.8 undertaking from PI confirming the same	
(a)	Identifying Information: Study Involves samples/data. If Yes, Specify Yes 🗖 No 🗖 NA 🗖	
	Anonymous/unidentified 🔲 Anonymized: Irreversibly Identifiable 🔲 reversibly coded 🔲 coded 💭	
	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.)	k
(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 I No Aybe I If yes, explain how you might use stored material/data in the future?	
	SECTION D: OTHER ISSUES	
10. PUB	SECTION D: OTHER ISSUES	
1 0. PUB (a)		
	LICATION, BENEFIT SHARING AND IPR ISSUES	
(a)	LICATION, BENEFIT SHARING AND IPR ISSUES Will the results of the study be reported and disseminated? If yes, specify. Yes No NA NA	
(a) (b)	LICATION, BENEFIT SHARING AND IPR ISSUES Will the results of the study be reported and disseminated? If yes, specify. Yes No NA NA Will you inform participants about the results of the study? Yes No NA NA Are there any arrangements for continued provision of the intervention for participants, if effective,	
(a) (b) (c)	LICATION, BENEFIT SHARING AND IPR ISSUES Will the results of the study be reported and disseminated? If yes, specify. Yes No NA Will you inform participants about the results of the study? Yes No NA Are there any arrangements for continued provision of the intervention for participants, if effective, once the study has finished? If yes describe in brief (Max 50 words) Yes No NA Is there any plan for post research benefit sharing with participants? If yes, specify	

elsewhere in the form? If yes, provide the details.

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	Version

Version 2.0 06

⁹For example, a data entry room, a protected computer etc.

SECTION E: DECLARATION AND CHECKLIST⁰

11. DE	1. 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 including responsible.				
	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.				
	If applicable, 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 (if required) 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 safety and confidentiality of study 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.				
	I/We have the following conflict of interest (PI/Co-PI):				
	1. 2.				
	I/We declare/confirm that all necessary government approvals will be obtained as per requirements wherever applicable.				
	Name of PI: Signature: Click here to enter a date.				

	Name of Co-PI: Signature: Click he	ere to e	enter a	date.			
	Name of Guide: Signature: Click he	re to e	nter a	date.			
	Name of HOD: Signature: Click her	e to ei	nter a c	late.			
12. Cl	IECKLIST	•	•	-			
S.No	Items	Yes	No	NA	Enclosure No.	EC Remai applicabl	-
ADMI	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 ¹¹						
					Ver	sion 2.0 08	

13.	Investigators Brochure drug/biologicals/device trials		oplicable	for					
14.	Participant Information Sheet(PIS) and Informed Consent Form (ICF)(English and translated)								
15.	Assent form for minors (12-18 years) (English and Translated)								
16.	Proforma/Questionnaire / Case Report Forms (CRF)/ Interview guides/ Guides for Focused Group Discussions (FGDs) (English and translated)								
17.	Advertisement/material to recruit participants (fliers, posters etc)								
PERMISSION FROM GOVERNING AUTHORITIES									
	Other Registration/ permissions	Required	Not required	Receiv	ved	Appli dd/m	ed ım/yy	EC Remark	S
18.	CTRI						date		
19.	DCGI					Enter	date		
20.	HMSC					Enter	date		
21.	NAC-SCRT					Enter	date		
22.	ICSCR					Enter	date		
23.	RCGM					Enter	date		
24.	GEAC					Enter			
25.	BARC					Enter			
26.	Tribal Board					Enter			
27.	Others (Specify)					Enter			
ANY OTHER RELEVANT INFORMATION/DOCUMENTS RELATED TO THE STUDY									
	Item		YES		NA	Enclo no.	sure	EC remarks	
28.									
29.									

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)

*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

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