

Application Form for Initial Review

अखिल भारतीय आयुर्वेद संस्थान

All INDIA INSTITUTE OF AYURVEDA (AIIA)

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

SECTION A - BASIC INFORMATION

1. (a) (b) (c)					
(d)	Department/	Division:		(e) Da	ate of Submission: Click here to enter a date.
(f)	Type of revie	w requested ¹ : om Review	Expedited Reviev	v 🗖	Full Committee Review
(g)	Title of the st	udy:			
	Acronym/ Sh	ort title, (If any):			
(h) (i)	Protocol num Details of Inv	· · · · · · · · · · · · · · · · · · ·		Versior	n number:
	Name	Designation and Qualification	Department and Institution	Address f	or communication ²
Pr	incipal Investiga	ntor/Guide			
Сс	o-investigator/st	udent/fellow			
j)		udies where applicar al Investigator at tim		ii) Co-	Investigator at time of submission:
(k)	Duration of t	he study:			

¹ Refer to National Ethical Guidelines for Biomedical & Health Research Involving Human Participants 2017on Page 36 Table 4.2. for the types of review

²Include telephone/mobile, fax numbers and email id

	at site	In Inc		Globally	🗖	
	Self-funding 📙	I	nstitutional funding L		nding agency 🔲	
	9	SECTION E	<mark>3 - RESEARCH REL</mark>	ATED INFO	ORMATION	
O\	ERVIEW OF RESEA	RCH				
(a)	Lay Summary of s	tudy ³ (withir	n 300 words)			
(b)	Type of study:					
	Basic Sciences		Clinical		Cross Sectional	
	Retrospective		Epidemiological/ Public Health	Ш	Case Control	
	Prospective		Socio-behavioral		Cohort	
	Qualitative		Dielerieel		Systematic Review	
	Quantitative		Biological samples/Data			
	Mixed Method		Any others (Specify)			
. M E	ETHODOLOGY Sample size/ No. o	-				
	Sample size/ No. of At site Control group	In Indi Study Grou ne sample siz	a Globally	In case of qu	ualitative study, menti	ion the criteria
(a)	Sample size/ No. of At site Control group Justification for the used for saturation Is there an extern	In Indi Study Grou ne sample siz on	a Globally p te chosen (100 words); y/ outsourcing involved	I for investig	_	
(a)	Sample size/ No. of At site Control group Justification for the used for saturation Is there an extern How was the scient	In Indi Study Grou ne sample siz on al laboratory ntific quality	a Globally p te chosen (100 words); // outsourcing involved of the study assessed?	I for investig	rations? ⁴ Yes	
(a)	Sample size/ No. of At site Control group Justification for the used for saturation Is there an extern	In Indi Study Grou ne sample siz on al laboratory ntific quality	a Globally p te chosen (100 words); y/ outsourcing involved of the study assessed? Review by	I for investig	_	□ NA□
(a)	Sample size/ No. of At site Control group Justification for the used for saturation Is there an extern How was the scien Independent extended.	In Indi Study Grou ne sample size on al laboratory ntific quality ternal	a Globally p te chosen (100 words); // outsourcing involved of the study assessed?	I for investig	rations? ⁴ Yes No Review within	□ NA□
(a)	Sample size/ No. of At site Control group Justification for the used for saturation Is there an extern How was the scient Independent exteriew	In Indi Study Grou ne sample size on al laboratory ntific quality cernal	a Globally p te chosen (100 words); // outsourcing involved of the study assessed? Review by Sponsor/Funder	I for investig	rations? ⁴ Yes No Review within	□ NA□
(a)	Sample size/ No. of At site Control group Justification for the used for saturation Is there an extern How was the scient Independent exteriew Review within more center research groups.	In Indi Study Grou ne sample size on al laboratory ntific quality cernal	a Globally p te chosen (100 words); // outsourcing involved of the study assessed? Review by Sponsor/Funder	I for investig	rations? ⁴ Yes	NA NA
(a)	Sample size/ No. of At site Control group Justification for the used for saturation Is there an extern How was the scient Independent exteriew Review within more center research group.	In Indi Study Grou ne sample size al laboratory ntific quality ternal nulti- group	a Globally p te chosen (100 words); // outsourcing involved of the study assessed? Review by Sponsor/Funder No Review	I for investig	rations? ⁴ Yes	□ NA□
(a)	Sample size/ No. of At site Control group Justification for the used for saturation Is there an extern How was the scient Independent exteriew Review within more center research group.	In Indi Study Grou ne sample size al laboratory ntific quality ternal nulti- group	a Globally p te chosen (100 words); // outsourcing involved of the study assessed? Review by Sponsor/Funder	I for investig	rations? ⁴ Yes	NA NA

4If participant samples are sent outside for investigations, provide details of the same and attach relevant documentation such as a etc.

SECTION C - PARTICIPANT RELATED INFORMATION

5. RECRUITMENT AND RESEARCH PARTICIPANTS (a) Type of participants in the study: Healthy Patient Vulnerable person/ Others volunteer Special groups (Specify) Who will do the recruitment? Participant recruitment methods used: Posters/ TV/Radio Patients / Telephone leaflets/Letters ads/social Family/Friends media/Institution visiting website hospitals Others (Specify) Yes No No NA (b) Will there be vulnerable person/special groups involved? ii. If yes, type of vulnerable person /special groups Children under 18 yrs Pregnant or lactating women Differently abled (Mental/Physical) Employees/Students/Nurses/ Staff Elderly Institutionalized Economically and socially disadvantaged Refugees/Migrants/Homeless Terminally III (stigmatized or rare diseases) Any other (Specify): Provide justification for inclusion/exclusion iii. Are there any additional safeguards to protect research participants? iv. Yes No No (c) Is there any reimbursement to the participant? If yes, Monetary non-monetary Provide details Yes No No Are there any incentives to the participant? If yes, Monetary non-monetary Provide details (e) Are there any participant recruitment fees/ incentives for the study provided to the PI/ Institution? Yes No 🗆 If ves. Monetary Non-monetary Provide details

	If yes, categorize th Less than Minima		Minimal risk	,	Yes No	
	Minor increase ov Low Risk ii. Describe the risk m	ver minimal risk or anagement strategy:	More than Mi	nimal Risk or Hig	h Risk	
(b)	What are the potential	benefits from the study	? Yes No	If yes, Direct	Indirect	
	For the participant					
	For the society/commu	ınity				
	For improvement in sci Please describe how th	ence e benefits justify the ris	ks			
(c)	Are Adverse Events exp	pected in the study ⁶ ?			Yes No No	NA 🔲
	Are reporting procedur	es and management str	ategies describe	d in the study?	Yes 🔲 No 🔲	
7. IN	NFORMED CONSENT					
7. In (a)		of consent? If yes, plea	se specify reasor	ns and skip to que	estion 8. Yes 🔲	No 🗖
(a) (b)	Are you seeking waiver Version number and da Version number and da	ate of Participant Inform ate of Informed Consent	ation Sheet (PIS)		estion 8. Yes 🗖	No 🗖
(a)	Are you seeking waiver Version number and da	ate of Participant Inform ate of Informed Consent ed for: Uerbal/ oral	ation Sheet (PIS) Form (ICF):	ed 🗖 /	Audio-Video	No 🗖
(a) (b)	Are you seeking waiver Version number and da Version number and da Type of consent planne Signed consent	ate of Participant Inform ate of Informed Consent ed for:	ation Sheet (PIS) Form (ICF):	ed		No 🗖
(a) (b)	Version number and da Version number and da Type of consent planner Signed consent Consent from LAR (If so, specify from	ate of Participant Informate of Informed Consent Verbal/ oral consent For children<7 yrs parental/LAR consent	ation Sheet (PIS) Form (ICF): Witnesse consent Verbal as from min 12 yrs) ale with pare	ed	Audio-Video (A/V) consent Written Assent from Minor (13- 18 yrs) along with	No 🗆
(a) (b) (c)	Are you seeking waiver Version number and day Version number and day Type of consent planner Signed consent Consent from LAR (If so, specify from whom)	ate of Participant Informate of Informed Consented for: Verbal/ oral consent For children<7 yrs parental/LAR consent	ation Sheet (PIS) Form (ICF): Witnesse consent Verbal as from min 12 yrs) ale with pare consent	ed	Audio-Video (A/V) consent Written Assent from Minor (13- 18 yrs) along with	No

⁶The term adverse events in this regard encompass both serious and non-serious adverse events.

(e)	Participant Informa English List the languages in	Local langu	ıage		ent Form (ICF other 🔲	(specify)			
(f)	If translation has no Provide details of C			viously	stored sampl	es if used in the study ⁷			
(g)	Elements contained in the Participant Information Sheet (PIS) and Informed Consent Form (ICF)								
	Simple language	Data shar	a/ Sample		Compensation	on for study related injury			
	Risks and discomforts		d to recontact		Statement th	hat consent is voluntary			
	Alternatives to participation	Con	fidentiality		Commerciali	ization/benefit sharing			
	Right to withdraw		age of		Statement th	hat study involves research			
	Benefits		rn of research		Use of photo	ographs/ identifying data			
	Purpose and procedure Others (Specify)	Payı	ment for icipation		Contact info Secretary of	rmation of PI and Member EC			
8. P (a	AYMENT/COMPENSA) Who will bear the PI				orocedures ⁸ ? onsor	Other agencies(specify)			
(b) Is there a provisio	n for free tre	eatment of rese	arch re	lated injuries	? Yes No E	NA 🗆		
(c	If yes, then who w	•		arch rel	ated SAE? If y	ves, specify. Yes 🔲 No 🎚	□ NA□		
	Sponsor 🔲 Ins	stitution/ Co	rpus funds 🔲	P	roject grants	Insurance			
(d) Is there any provis					ne relatedness is determine Yes 🔲 No	d for		
(e) Is there a provision specify.	for ancillary	care for unrela	ted illn	ess during the	e study period? If yes, pleas Yes No	e NA		
⁷ In 1	formation on re-consent requi	irements can be f	ound at National Ethi	cal Guide	lines for Biomedica	ıl & Health Research Involving Human			

Participants 2017, Page 54 in Section 5.8

⁸Enclose undertaking from PI confirming the same

9. ST	DRAGE AND CONFIDENTIALITY
(a)	Identifying Information: Study Involves samples/data. If Yes, Specify Yes No NA 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.)
(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? If yes, explain how you might use stored material/data in the future?
	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 NA
(b)	Will you inform participants about the results of the study?
(c)	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 \square No \square NA \square
(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, Please provide details Yes No NA NA
(f)	Do you have any additional information to add in support of the application, which is not included elsewhere in the form? If yes, provide the details. Yes \square No \square
⁹ For	example, a data entry room, a protected computer etc.

Version 2.0 06

SECTION E: DECLARATION AND CHECKLIST 10

11. D	ECLARATION (Please	e tick as applica	able)							
	I/We certify that t	he 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 institutions where	•	_	delines of the institute and affiliated/collaborating ed.						
	I/We will ensure will adhere to the	•		this study are qualified, appropriately trained and ved protocol.						
	I/We declare that	the expenditur	re in case of	injury related to the study will be taken care of.						
	If applicable, I/We provided, if applic		an undertaki	ng of what will be done with the leftover samples is						
	I/We confirm that	t we shall subm rotocols, progr		ocol amendments, adverse events report, significant if required) and a final report and also participate in						
	I/We confirm that	we will mainta	ain accurate	and complete records of all aspects of the study.						
	I/We will protect and biological sam		participants	and assure safety and confidentiality of study data						
				gators, researchers and/or close relative(s), have no with the sponsor(s) and outcome of study.						
	I/We have the foll	lowing conflict	of interest (I	PI/Co-PI):						
	1. 2.									
	I/We declare/cor requirements who		•	government approvals will be obtained as per						
	Name of PI:	Signature:		Click here to enter a date.						
	Name of Co-PI:	Signature:		Click here to enter a date.						

	Name of Guide: Name of HOD:	Signature: Signature:	Click he						
12. CF	HECKLIST	Items		Yes	No	NA	Enclosure No.	EC Remarks applicable)	(If
ADMI	NISTRATIVE REQUIREM	ENTS							
1.	Cover letter								
2.	Brief CV of all Investiga	ators							
3.	Good Clinical Practice last 3 years	(GCP) training	of investigators in						
4.	Approval of Scientific (Committee							
5.	EC clearance of other	centers*							
6.	Agreement between c	ollaborating par	tners*						
7.	MTA between collabor	rating partners*							
8.	Insurance policy/certif	icate							
9.	Evidence of external la externally outsource certification								
10.	Copy of contract or agor donor agency	greement signed	d with the sponsor						
11.	Provide all significan leading to a negative other ECs/Regulatory (whether in same modification(s) to prof	decision or mo authorities fo location or	dified protocol) by r proposed study						
PROP	OSAL RELATED								
12.	Copy of the detailed p	rotocol ¹¹							
13.	Investigators Broc drug/biologicals/devic	chure (If e trials)	applicable for						
14.	Participant Informati	on Sheet (PI	S) and Informed						

	Consent Form (ICF)(English a	nd translate	ed)						
4.5					_				
15.	Assent form for minors (Translated)	12-18 year	rs) (English	and L	_	Ш	Ш		
16.	Proforma/Questionnaire / Interview guides/ Guides for (FGDs) (English and translate	-							
17.	Advertisement/material to posters etc)	recruit par	rticipants (fliers,					
PERM	IISSION FROM GOVERNING AU	JTHORITIES							
	Other Registration/ permissions	Required	Not required	Receive	d	Applie dd/mr		EC Remark	s
18.	CTRI					Enter			
19.	DCGI					Enter	date		
20.	HMSC					Enter	date		
21.	NAC-SCRT					Enter	date		
22.	ICSCR					Enter	date		
23.	RCGM					Enter	date		
24.	GEAC					Enter	date		
25.	BARC					Enter	date		
26.	Tribal Board					Enter	date		
27.	Others (Specify)					Enter	date		
ANY (OTHER RELEVANT INFORMATION	ON/DOCUM	IENTS RELA	TED TO T	HE S	STUDY			
	Item		YES	NO N	Α	Enclos no.	ure	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)