

### (Annexure 15)

## **Initial Review Form for Multicentric Research**

# अखिल भारतीय आयुर्वेद संस्थान All INDIA INSTITUTE OF AYURVEDA (AIIA)

**EC Ref. No.** (for office use):

PART 1 (To be filled by coordinating PI)

#### **SECTION A - BASIC INFORMATION**

General Instructions: a) Tick one or more as applicable. Mark NA if not applicable. Attach additional sheets if required b) For submission to Designated Ethics Committee and to be shared with PIs at Participating Centres

1. (a)	Name of Institute under which Designated Ethics Committee is constituted:
(b) (c)	Name of the Ethics Committee: Name of Coordinating Principal Investigator:
(d)	Designation and Qualification:
(e)	Department/Division: (e) Date of Submission: Click here to enter a date.
(f)	Address for communication (include email and mobile no.)
(g)	Type of review requested¹:  Exemption from Review  Expedited Review  Full Committee Review
(h)	Title of the study:  Acronym/ Short title, (If any):
(i)	Protocol number (If any): Version number: Date: Click here to enter a date.
(j)	Number of studies where applicant is a:  i) Principal Investigator at time of submission:  ii) Co-Investigator at time of submission:
(k)	Duration of the study:
2.	FUNDING DETAILS AND BUDGET
(a)	Total estimated budget for study: At site In India Globally
(b)	Self-funding Institutional funding Funding agency (Specify)

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

## **SECTION B – RESEARCH-RELATED INFORMATION**

3.	OV (a)	ERVIEW OF RESEA Lay Summary of s		n 300 wo	rds)				
	(b)	Type of study: Basic Sciences Retrospective Prospective Qualitative Quantitative Mixed Method		Health Socio-be Biologic samples			Cross Sec Case Con Cohort Systemat		
4.	ME	THODOLOGY							
	(a)	Sample size/ No. At site Control group Justification for the used for selection	In Indi Study Grou he sample siz	р	Globally	n case of qua	litative st	udy, mentic	on the criteria
	(b) (c)	Is there an extern How was the scie Independent ext review	entific quality	of the st			Revi	No No lew within institution	
		Review within m centre research	ļ	□ N	o Review				
		Date of review: Comments of Sci	ientific Comm	nittee, if	any (100 words	5)		Click here to (	enter a date.

<sup>&</sup>lt;sup>2</sup>Summarize in the simplest possible way such that a person with no prior knowledge of the subject can easily understand it.

<sup>3</sup>If participant samples are sent outside for investigations, provide details of the same and attach relevant documentation such as an MTA/ MoU etc.

#### **SECTION C - PARTICIPANT RELATED INFORMATION**

#### 5. RECRUITMENT AND RESEARCH PARTICIPANTS (a) Type of participants in the study: Others Healthy Patient Vulnerable person/ volunteer Special groups (Specify) Who will do the recruitment? Participant recruitment methods used: TV/Radio Telephone Posters/ Patients / leaflets/Letters Family/Friends ads/social media/Institution visiting website hospitals Others (Specify) Yes No No NA (b) Will there be vulnerable person/special groups involved? If yes, type of vulnerable person /special groups ii. 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): iii. Provide justification for inclusion/exclusion iv. Are there any additional safeguards to protect research participants? Yes No No (c) Is there any reimbursement to the participant? If yes, Monetary non-monetary Provide details (d) Are there any incentives to the participant? Yes No No 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 yes, Monetary Non-Monetary Provide details

	i. Are there any antici	oated physical/social/p	sycho	logical dis	comforts,	/ risk to	participants? Yes No No	
	If yes, categorize the Less than Minimal		Min	imal risk				
	Minor increase ove		Moi	re than M	inimal Ris	sk or Hi	gh Risk	
	ii. Describe the risk ma	nagement strategy:						
(b)	What are the potential b	enefits from the study	? Ye	es No	If yes,	Direc	t Indirec	t
	For the participant							
	For the society/commun	ity						
	For improvement in scie	nce						
	Please describe how the	benefits justify the ris	ks					
(c)	Are Adverse Events expe	ected in the study <sup>5</sup> ?					Yes No 🗖	NA 🗖
	Are reporting procedure If Yes, Specify	•	ategie	s describe	ed in the s	study?	Yes 🔲 No 🔲	
7. II	NFORMED CONSENT							
(a)	Are you seeking waiver of	of consent? If yes, plea	se spe	cify reaso	ns and sk	ip to q	uestion 8. Yes 🗖	No
(b)	Version number and dat	e of Participant Inform	ation :	Sheet (PIS	5):			- <del>-</del>
	Version number and dat	e of Informed Consent			s):			
(b)	Version number and dat Type of consent planned	e of Informed Consent I for:	Form	(ICF):			Audio Vidoo	
	Version number and dat	e of Informed Consent I for:	Form			•	Audio-Video (A/V) consent	
	Version number and dat Type of consent planned	e of Informed Consent I for:   Verbal/ oral   consent	Form	(ICF): Witness	ssent nor (7- along			
(c)	Version number and dat Type of consent planned Signed consent  Consent from LAR (If so, specify from whom)  Other (specify)	e of Informed Consent I for:  Verbal/ oral consent For children<7 yrs. parental/LAR consent	Form	Witness consent Verbal a from mii 12 yrs.) a with par	ssent nor (7- along		(A/V) consent Written Assent from Minor (13- 18 yrs) along with	
	Version number and dat Type of consent planned Signed consent  Consent from LAR (If so, specify from whom)	e of Informed Consent I for:  Verbal/ oral consent For children<7 yrs. parental/LAR consent	Form	Witness consent Verbal a from mir 12 yrs.) a with par consent	ssent nor (7- along		(A/V) consent Written Assent from Minor (13- 18 yrs) along with	
(c)	Version number and date Type of consent planned Signed consent  Consent from LAR (If so, specify from whom)  Other (specify) Who will obtain the info	e of Informed Consent I for:    Verbal/ oral     consent   For children<7 yrs.     parental/LAR     consent	Form	Witness consent Verbal a from mir 12 yrs.) a with par consent	ssent nor (7- along ental		(A/V) consent Written Assent from Minor (13- 18 yrs) along with parental consent	
(c)	Version number and dat Type of consent planned Signed consent  Consent from LAR (If so, specify from whom)  Other (specify)  Who will obtain the info PI/Co-I	e of Informed Consent I for:    Verbal/ oral     consent   For children<7 yrs.     parental/LAR     consent	Form	Witness consent Verbal a from mir 12 yrs.) a with par consent	ssent nor (7- along ental		(A/V) consent Written Assent from Minor (13- 18 yrs) along with parental consent	
(c)	Version number and dat Type of consent planned Signed consent  Consent from LAR (If so, specify from whom)  Other (specify)  Who will obtain the info PI/Co-I	e of Informed Consent I for:    Verbal/ oral     consent   For children<7 yrs.     parental/LAR     consent	Form	Witness consent Verbal a from mir 12 yrs.) a with par consent	ssent nor (7- along ental		(A/V) consent Written Assent from Minor (13- 18 yrs) along with parental consent	

<sup>&</sup>lt;sup>4</sup>For categories of risk refer to National Ethical Guidelines for Biomedical & Health Research Involving Human Participants 2017. Page 6 in Table 2.1

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

(e)	English 🗖	Loca	neet (PIS) and Informo language h translations were d		sent Form (ICF other 🗖	(specify)	
(f)			n done, please justify requirement for pre	viously	stored sample	es if used in the study <sup>6</sup>	
(g)	Elements contained	d in the	Participant Informat	ion Sh	eet (PIS) and Ir	nformed Consent Form (ICF	:)
	Simple language		Data/ Sample sharing		Compensatio	on for study related injury	
	Risks and discomforts		Need to recontact		Statement th	nat consent is voluntary	
	Alternatives to participation		Confidentiality		Commerciali	zation/benefit sharing	
	Right to withdraw		Storage of samples		Statement th	nat study involves research	
	Benefits		return of research results		Use of photo	graphs/ identifying data	
	Purpose and procedure Others (Specify)		Payment for participation		Contact infor Secretary of	rmation of PI and Member EC	
<b>8.</b> P. (a)	AYMENT/COMPENS. Who will bear the	costs	related to participation		procedures <sup>7</sup> ? onsor	Other agencies(specify)	
(b)	Is there a provisio	n for f	ree treatment of rese	earch re	elated injuries?	Yes No E	NA 🗖
(c)	•	•	vide the treatment? ompensation of resea	arch re	lated SAE? If y	es, specify. Yes 🗖 No 🗓	□ NA□
	Sponsor 🔲 In:	stitutio	n/ Corpus funds	] F	Project grants	Insurance 🗖	
(d)			r medical treatment of during the study per		•	ne relatedness is determine Yes No	
(e)	Is there a provision specify.	for an	cillary care for unrela	ted illr	ess during the	study period? If yes, pleas Yes No NA	
-	rmation on re-consent requi Participants 2017, Page 5- lose undertaking from PI col	4 in Sectio	on 5.8	ical Guide	lines for Biomedical	& Health Research Involving Human	

9. ST	ORAGE AND CONFIDENTIALITY			
(a)	Identifying Information: Study Involves samples/data. If Yes, Specify	Yes	No 🗖	NA 🔲
	Anonymous/unidentified Anonymized: Irreversibly reversibly coded coded	Identif	iable	
	If identifiers must be retained, what additional precautions will be taken to / data is safeguarded? (e.g. data stored in a cabinet, password protected co			ss is limited
(b)	Who will be maintaining the data pertaining to the study?			
(c)	Where will the data be analyzed <sup>7</sup> 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?	Yes	No 🗖	Maybe 🗖
	SECTION D: OTHER ISSUES			
10. PUB	LICATION, 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 once the study has finished? If yes describe in brief (Max 50 words)	participar Yes 🔲 N		
(d)	Is there any plan for post research benefit sharing with participants? If yes, s	specify Yes 🗖	No 🗖	na 🗖
(e)	Is there is any commercial value or a plan to patent/IPR issues. If yes, Please	e provide o Yes 🔲		NA 🔲
(f)	Do you have any additional information to add in support of the application, elsewhere in the form? If yes, provide the details.	, which is Yes		uded
<sup>7</sup> Foi	r example, a data entry room, a protected computer etc.			

# **SECTION E: CHECKLIST FOR COORDINATING PI**

11. CH	IECKLIST								
S.No	Ito	ems			Yes	No	NA	Enclosure No.	EC Remarks
ADMI	NISTRATIVE REQUIREMENT	ΓS							
1.	Cover letter								
2.	Brief CV of all Investigator								
3.	Good Clinical Practice (Go in last 3 years	CP) training	of investig	ators					
4.	Approval of Scientific Advisory Committee/ Any		' NTF/ Ce	entral					
5.	Agreement/MTA / LOA be partners		borating						
6.	Insurance policy/certificat	e							
7.	Evidence of external laboran externally outsourced certification	•							
8.	Copy of contract or ag sponsor or donor agency	reement s	igned with	the					
PROP	OSAL RELATED								
9.	Copy of the detailed proto	col							
10.	Participant Information Consent Form (ICF) (Englis			rmed					
11.	Assent form for minors Translated)	(12-18 yea	rs) (English	and					
12.	Proforma/Questionnaire , Interview guides/ Guid Discussions (FGDs) (Englis	des for F	ocused G	CRF)/ Group					
13.	Advertisement/material to posters etc.)	o recruit pa	rticipants (1	fliers,					
PERM	ISSION FROM GOVERNING	AUTHORIT	IES					-	
	Other Registration/ permissions	Required	Not required	Recei	ived	Appli dd/m	ed m/yy	EC Remark	s
14.	CTRI <sup>8</sup>					Enter			
15.	HMSC <sup>9</sup>					Enter	date		
16.	Tribal Board					Enter	date		
17.	Any Other					Enter	date		
ANY C	THER RELEVANT INFORMA	ATION/DOC	UMENTS RI	ELATE	тот	HE STU	IDY		
	Item		YES	NO	NA	Enclo no.	sure	EC remarks	
18.									

 ${\it ^8CTRI: Clinical\ Trial\ Registry-\ India,\ ^9HMSC:\ Health\ Ministry's\ Screening\ Committee}$ 

## PART 2 (To be filled by S-PI at the Participating Centre)

General Instructions: a) Tick one or more as applicable. Mark NA if not applicable. Attach additional sheets if required b) For submission to Participating Ethics Committee (PEC) and to be shared with coordinating PI

### **SECTION A - BASIC INFORMATION**

1. a) b) c) d) f)	ADMINISTRATIVE DETAILS  Name of the institute under which PEC is constituted:  Name of the Ethics Committee:  Name of Site Principal Investigator:  Designation/ Qualification:  Address for communication (include mobile no. and email address):
g)	Expected duration of the study: Estimated budget at the participating site:
	SECTION B - RESEARCH INFORMATION
<b>1.</b> a)	OVERVIEW OF RESEARCH Briefly describe the role of the participating center in the study (50-100 words):
b)	Briefly mention local changes made in protocol, if any:
c)	Type of review requested:  Exemption from Review
	SECTION C – PARTICPANT RELATED INFORMATION
1. a) b)	PATIENT RECRUITMENT AND RESEARCH PATIENTS  Number of participants to be recruited at site:  Site specific/ community concerns, if any
c)	Briefly mention local changes in Recruitment/ Advocacy material:
d)	Copy of the Local Recruitment/ Advocacy material: Yes 🔲 No 🔲
<b>2.</b> a)	INFORMED CONSENT Who will obtain the informed consent?  S-PI/Co-S-PI Nurse/Counselor Research Staff Other (Specify)
	Any tools to be used
b) c) d) e)	Language/s ICD is translated in:  Version number and date of the Participant Informed Sheet (PIS):  Version number and date of the Informed Consent form (ICF):  Copy of the Local ICD translations enclosed: Yes No

f)	Back translation of the ICD in English with the translation certificate Yes $lacksquare$ No $lacksquare$
g)	Changes made in informed consent form (ICF), if any:
h)	Copy of the audio / visual transcript for consent enclosed, if any: Yes No
3.	DATA AND STORAGE
i)	Brief details on data collection, storage, sharing, transfer, if any?
	SECTION D – OTHER ISSUES
a)	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 $\square$
	to a notation and the second and seconds.

## SECTION E – CHECKLIST FOR S-PI AT PARTICPATING CENTER

1. CHEC	1. CHECKLIST							
Sr.No	Items	Yes	No	NA	EnclosureNo.	EC Remarks		
ADMINI	STRATIVE REQUIREMENTS							
1.	Cover letter							
2.	Brief CV of Site Principal Investigator / other site Co-PI							
3.	Good Clinical Practice (GCP) training of investigator in last 3 years							
4.	Agreement between collaborating partners							
5.	MTA between collaborating partners							
6.	Insurance policy/certificate							
PROPOS	SAL RELATED							
7.	Copy of the modified protocol							
8.	Participant Information Sheet (PIS) and Informed Consent Form (ICF) (English and translated)							
9.	Assent form for minors (12-18 years) (English and Translated)							
10.	Proforma/Questionnaire / Case Report Forms (CRF)/ Interview guides/ Guides for Focused Group Discussions (FGDs) (English and translated)							
11.	Advertisement/material to recruit participants (fliers, posters etc.)							
12.	Any other relevant information/documents related to the study							