

## (Annexure 8) Application form for Clinical Trials अखिल भारतीय आयुर्वेद संस्थान

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

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

Title of study:							
Principal Investigator (Name, Designation and Affiliation):							
1.	Type of clinical trial		Regulatory trial Academic trial				
	CTRI registration number:		NABH accreditation number EC registration number:				
2.	If regulatory trial, provide stat	ovide status of CDSCO permission letter					
	Approved and letter attached $\square$						
	Applied, under process						
	Not applied (State reason)						
3.	Tick all categories that apply to	o your					
	Phase - I		Phase II				
	Phase III		Phase IV or Post Marketing Surveillance				
	Investigational medicinal products		Investigational New drug				
	Medical devices		New innovative procedure				
	Drug/device combination		Bioavailability/Bioequivalence studies				
	Non-drug intervention		Repurposing an existing intervention				
	Indian system of medicine (AYUSH)		Stem cells				
	Phytopharmaceutical drug		Approved drug for any new indication or new route of administration				
	Others (specify)						
4.	Trial design of the study (May	choos	e more than one)				
	I. Randomized Nonrandomized Parallel Cross-over		Factorial Stratified Adaptive Comparison trial				
	Cluster		Superiority trial				
	Matched pair		Non-inferiority trial				

	Others (specify)		Equivalence trial	
	II. If there is randomiza group(s)?	ition, how will the	e participants be allocated to the	control and study
	III. Describe the method of	of allocation concea	alment (blinding / masking), if applica	ble
5.	List the primary / secondary	outcomes of the t	rial.	
6.	Is there a Contract Research Agency such as public relati If yes, Name and Contact d	on/Human resourc	D) /Site Management Organization (S e? Yes	MO) / Any Other No
	State how the CRO/SMO/ag	gency will be involve	ed in the conduct of the trial (tick all	that apply)
	Project management		Clinical and medical monitoring	
	Regulatory affairs		Data management	
	Statistical support		Medical writing	
	Site management		Audits, quality control, quality assurance	
	Finance management		Recruitment and training	
	Administrative support		Others (specify)	
7.	Please provide the following	g details about the	intervention being used in the proto	col
	I. Drug/s, device/s and/or b	iologics; If yes, prov	vide regulatory approval details Yes	No□ NA□
	II. Already approved drugs dosage form / route of adm		of two or more drugs with new indi provide details Yes	
	III. Provide contact details biologics	of who prepared	d and /or is manufacturing the dr	ug/s, device/s and
	IV. Provide details of patent	of the drug/s, devi	ice/s and biologics.	

8.	Describe in brief any preparatory work or site preparedness for the protocol? Yes No NA						
	If yes, (100words)						
9.	Is there an initial scr	reening/ use of existing database for participant selection?	Yes No NA NA				
	If Yes, provide detai	ls <sup>22</sup>					
10.	•	any anticipated incidence, frequency and duration of adverse events related to the n? If yes, provide details of arrangements made to address them. Yes No NA					
11.	Does the study use a	a placebo? e of the placebo and risks entailed to participants.	Yes No NA				
12.	Will current standard of care be provided to the control arm in the study?  Yes No NA NA If no, please justify.						
13.	Are there any plans to withdraw standard therapy during the study ?If yes, please justify.  Yes No No NA						
14.	Are there any rules to stop the protocol in case of any adverse events? If yes, please specify. Yes $\square$ No $\square$ NA $\square$						
15.	Does the study have	e a Data and Safety Monitoring Plan? If no, please justify.	Yes No				
16.	Participant Informat	tion Sheet (PIS) and Informed Consent Form (ICF)					
	English 🔲	Local language  (Certified that local version (s) is/are a true translation of the English version and can be easily understood by the participants)					
	List the languages in which translations were done						
	Justify if translation not done						
	<sup>22</sup> In order to select participants for your protocol does the protocol require you to screen an initial population or refer to an existing database before shortlisting participants. If yes, provide details on the same						

Version 2.0 03

17. 18.	Involvement/conso	Yes No NA NA Yes No No					
	i. Is the PI registered with Medical Council of India (MCI) or the State Medical Council registration?  Please provide details.  Yes No						
	ii. Is the PI trained	in GCP in last 3 years? If yes, please enclose certificate	Yes No 🗆				
Sig	nature of PI:	Click here to enter a date.					