(Annexure 1)



Application Form for Expedited Review

* Eluru -534005	(Name of the Institution) EC Ref. No.* (For office us	se):
	ame, Designation and Affiliation):	
	sons why expedited review from EC is requested 12 ?	
	es non-identifiable specimen and human tissue from sources like blood banks, tissue bank er clinical samples.	ks and
	es clinical documentation materials that are non-identifiable (data, documents, records).	
iii. Modific	cation or amendment to approved protocol (administrative changes/correction of typogra	aphical \square
iv. Revise	d proposal previously approved through expedited review, full review or continuing revie ed proposal.	ew of \square
v. Minor d	deviation from originally approved research causing no risk or minimal risk.	
_	ss/annual report where there is no additional risk, for example activity limited to data ana ted review of SAEs/unexpected AEs will be conducted by SAE subcommittee.	alysis. \square
vii. For mu	Iticentre research where a designated EC has approved the proposal, a participating EC	may \square
	participating centre specific information and modifications in the study proposal thro	_
	c to the centre.	ues ilivolveu
-	rch during emergencies and disasters (See Section 12 of ICMR Ethical Guidelines, 2017).	
ix. Any o	other (please specify)	
2. Is waiver of 0	consent being requested?	Yes 🗆 No 🗆
3. Does the res	search involve vulnerable persons ¹³ ?	Yes □ No □
If Yes give d	details:	
·····		
Signature o	of PI:dd	mm yy
Comments	of EC Secretariat:	
Signature of	f Member Secretary:	mm yy

 ¹² Refer to National Ethical Guidelines for Biomedical & Health Research Involving Human Participants 2017, Page 51 Table 4.2
¹³ For details, refer to application for initial review, Section-C, 5(b)
* In case this is first submission, leave it blank