

## **ETHICAL REVIEW COMMITTEE** University of Health Sciences, Lahore

Khayaban-e-Jamia Punjab, Lahore Office: 042-99231304-09, Fax 042-99231857

Title of the Project												
Name of Researcher												
Discipline/Specify Specialty		Ph.D 🔲	M.Phil 🗖	MD 🗖	ms 🗖	mds 🗖	мнре 🗖	M.Sc. 🗖	Resea	rch proj	ect 🔲	
Supervisor												
Institution												
Study Design												
Study Population												
	Please ans	wer each	question by	ticking th	ne approp	riate box				YES	NO	N/A
1.	Have you, l	oefore filli	ing in this for	m, read a	a relevant	research	ethics guide	eline of anim	nal,			
	human or b	biological material research?										
2.		ect multidisciplinary with involvement of different departments / institutes?										
3.			f the project	-				riate scientif	ic			
		experience for execution of the project under consideration?										
		study involve participants who are particularly vulnerable? (e.g. refugees,										
4.	-	, victims of violence, patients with sensitive medical conditions e.g. HIV/AIDS,										
	-		e unable to g	give infor	med cons	ent (e.g. c	hildren, pe	ople with				
	learning dis											
5.		•	ve archival m			-			ý			
	-	nd the material cannot be traced to its origin or cannot be accessed?										
6.		wer to the items 2,4,5 is yes, then have you taken formal approval from the ting institutes (organizations (centers?							2			
		ating institutes/organizations/centers? study involve discussion of sensitive topics (e.g. sexual activity, drug use) or										
7.		nbarrassment, psychological stress or anxiety or cause harm or negative										
7.	consequen											
		any potential conflict of interest relating to the study?										
8.		in have you declared the nature of conflict in the prescribed proforma? (This										
0.		on may be made a part of any existing proforma)										
9.			ments (other				nd comper	sation for ti	me)			
	be offered	red to participants?										
10.	Will drugs,	placebos,	or other sub	ostances (	e.g. food	or drink c	onstituents	, dietary				
	supplemen	plements) be administered to the study participants?										
11.			es, then is th	• •		-	red with Dr	ug Regulato	ry			
		ity of Pakistan according to DRAP Act 2012?										
	-	es available online at ww.dra.gov.pk/userfiles1/file/ProcedureforClinicalTrialApplications.pdf)										
		-	•								<u>                                     </u>	
12.		-	ve handling,	-		-	of Infectiou	s agents, to>	kins,			
			genic to hum								↓ ↓	
13.	If answer to	o 12 is yes	s, then are th	e standa	rd biosafe	ety measu	res (contam	ination cont	trol,			



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	spill response, waste management, use of protective apparel, and inventory control)		
	ensured for the execution of this project?		
14.	If the study involves distribution of questionnaires to the participants, has the right to		
	Response Omission and Anonymity been provided to them?		
15.	In case of animal studies, have you considered alternatives (in vitro systems, computer		
	simulations and/or mathematical models) to reduce or replace the use of animals as far		
	as possible?		
16.	Will you make sure that the health of humans/animals be given prior consideration and		
	avoid or minimize discomfort, distress, and all procedures will be kept aseptic, painless		
	and minimally intrusive?		
17.	Will you provide adequate care to all humans/animals and ailing study subjects shall be		
	properly treated by the qualified care providers and will be removed from further study?		
18.	Will you ensure confidentiality and data protection related to study participants?		
10.			
19.	Will you share study findings with the participants and ERC when asked?		
20.	Will you make sure that the results are only used for research purpose and information		
	disseminated only through research publication / conference papers/presentations?		
21.	Have you attached the consent form with explicit right of the participants to withdraw		
	from study at their will?		

**NOTE:** Duly filled and signed Proforma should be submitted with the Registration Department, along with the following documents:

- I. Title Page of Research Synopsis
- II. SRC (Synopsis Review Committee) Approval Letter
- III. Project Summary
- IV. Methodology/Research Proformas, if any
- V. Statistical Analysis
- VI. Informed consent programme designed for the study subjects, if applicable
- VII. Consent from the collaborating institution, if applicable (attach the letter of approval)

A Copy of the filled Proforma may also be emailed (Microsoft Word as well as PDF file) at **esynopsis@uhs.edu.pk** 

**Declaration:** I/We declare that all the information given in this form and written in the proposal is correct and I/We will abide by the ethical guidelines relevant to this research. I/We will reapply for ethical approval if there is a significant change or revision in the design or protocol of the proposed study.

Researcher	Supervisor	Co-Supervisor(s)	Head of Department
<b>Signature</b> Name: Cell #	<b>Signature</b> Name: Designation:	<b>Signature</b> Name: Designation:	<b>Signature</b> Name: Department:
	Department: Institution	Department: Institution	