

INSTITUTIONAL ETHICS COMMITTEE (IEC)

Seth G. S. Medical College and KEM Hospital, Mumbai.

Annexure 2

AX 02/ SOP 20/V 7

Checklist-Requirements for Research Involving Pregnant Women & Fetuses

-	HIS RESEARCH INVOLVES PREGNANT WOMEN OR FETUSES						
No.		Yes	No	NA			
1.	Where scientifically appropriate, preclinical studies, including studies on pregnant						
	animals, and clinical studies, including studies on non-pregnant women, have been						
	conducted and provide data for assessing potential risks to pregnant women and						
	fetuses;						
2.	The risk to the fetus is not greater than minimal, or any risk to the fetus which is						
	greater than minimal is caused solely by interventions or procedures that hold out						
	the prospect of direct benefit for the woman or the fetus;						
3.	Any risk is the least possible for achieving the objectives of the research;						
4.	The woman's consent or the consent of her legally authorized representative is						
	obtained in accord with the informed consent provisions, unless altered or waived.						
5.	The woman or her legally authorized representative, as appropriate, is fully informed						
	regarding the reasonably foreseeable impact of the research on the fetus.						
6.	No inducements, monetary or otherwise, will be offered to terminate a pregnancy;						
7.	Women's participation in the research will not have an effect on the decisions by						
	investigator with respect to the timing, method or procedures used to terminate a						
	pregnancy; and						
8.	The decision of investigator determining the viability of a fetus will not have an						
	effect if the women participates in the research.						
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A.	Fetuses of uncertain viability	Yes	No	NA
1.	Does the research hold out the prospect of enhancing the probability of survival of			
	the particular fetus to the point of viability, and any risk is the least possible for			
	achieving the objectives of the research ;			
	OR	ı	1	.4
	The purpose of the research is the development of important biomedical knowledge			
	which cannot be obtained by other means and there will be no risk to the fetus			
	resulting from the research ;			
2.	The legally effective informed consent of either parent of the fetus or, if neither			
	parent is able to consent because of unavailability, incompetence, or temporary			
	incapacity, the legally effective informed consent of either parent's legally			
	authorized representative is obtained.			
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В.	Nonviable fetuses	Yes	No	NA
1.	Vital functions of the fetus will not be artificially maintained;			
2.	There will be no risk to the fetus resulting from the research;			
3.	The purpose of the research is the development of important biomedical knowledge			
	that cannot be obtained by other means; and			
4.	The legally effective informed consent of both parents of the fetus will be obtained			
	except that the waiver and alteration provisions do not apply. However, if either			
	parent is unable to consent because of unavailability, incompetence, or temporary			
	incapacity, the informed consent of one parent of a nonviable fetus will suffice to meet			
	the requirements of this paragraph. The consent of a legally authorized representative			
	of either or both of the parents of a nonviable fetus will not suffice to meet the			
	requirements of this paragraph.			
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9	Signature of Principal Investigator: Date:			
	IEC Office use only			
Comr	ments			
	ary Reviewer Signature & Date:			