Annexure 5 AX 05/SOP 05/V6



Date:

Delegation of Responsibilities of Study team

INSTITUTIONAL ETHICS COMMITTEE (IEC)

Seth GS Medical College and KEM Hospital, Mumbai.

Drojoct	Pogistration No	
FIUICUL	i vedisti ation i vo	

ud y Title		
Name	Role	No.
	Principal Investigator	1
	Co-Investigator	2
	Co-Investigator	3
	Co-investigator	4
	Co-Investigator	5
	Co-investigator	6
	Study co-ordinator *	7
	Study co-ordinator *	7
	Laboratory Technician	8
		9
		10

(Please place tick marks against assigned duties for each member in the following table)

Code	TASKS	Role									
		1	2	3	4	5	6	7	8	9	10

^{*} Study coordinator may preferably be a person specifically appointed for coordinating the clinical trial; other than the staff member (assistant / associate professor)

A	All relevant documents pertaining to protect blinding					
В	Research participants selection/ Screening					
С	Obtain informed consent					
D	Evaluate inclusion/ exclusion criteria					
E	Conduct the visit assessments					
F	Physical examination					
G	Complete the source documents					
Н	Complete Case Record Form					
I	Final review and sign Case Record Form					
J	Collect laboratory safety test samples					
K	Processing of blood samples					
L	Preparing aliquots & keeping a track of the samples sent					
М	Review & sign of the lab repots					
N	Receive the study drug, , document drug dispensing, storage & accountability					
0	Person to whom research participants should contact in					

	case of adverse event					
Р	Report all serious adverse events					
Q	Follow up of Serious Adverse Event					
R	Maintaining study site master file					
S	In-charge of inventory & supplies					
Т	Archiving of study documents					
U	Resolution of queries					
V	Overall coordination and supervision					

Signature with dat	e of Principal Investigator:	
Jigijaluje Wilii ual	e di Fillicipal lilvestigator.	