Annexure 11 Participant Informed Consent Form

Protocol Study number:	
Patient identification number for this study:	
Title of the project:	
Name of Principal investigator:	Mobile No
explained in detail to me, in a language tha that I have had the opportunity to ask questi The nature and purpose of the study and it other relevant details of the study have be voluntary and that I am free to withdraw f medical care or legal right being affected. I understand that the information collected of my medical notes may be looked at by individuals to have access to my records. I a	ts potential risks / benefits and expected duration of the study, and en explained to me in detail. I understand that my participation is from the study at any time, without giving any reason, without my about me from my participation in this research and sections of any y responsible individuals from PIMS. I give permission for these agree to take part in the above study.
(Signatures / Left Thumb Impression of H	
	Son/Daughter/spouse of:
This is to certify that the above consent has	been obtained in my presence
This is to certify that the above consent has	been obtained in my presence.
Place	Date:
Signatures of the Principal Investigator	
1) Witness – 1	2) Witness – 2
Signature	Signature
Name:	Name:
■	Address: ch for (1) Patient (2) Researcher (3) Institution translation in simple understandable Hindi on their own)