Annexure 10

Participant Information Sheet (PIS)

The project must be accompanied by the Participant information sheet addressed to the patient or participant or parent/ guardian, in case of minor. While formulating the participant information sheet, the investigator must provide the subjects with the following information in English and Hindi, in a simple layman's language which can be understood by them, in a narrative form, directed to the participant/ LAR, covering all the points:

- 1. Study Title
- 2. Aims and methods of the research study
- 3. Expected duration of participation
- 4. The benefits to be expected from the research to the participant or to others
- 5. Any risk or discomfort to the participant associated with the study
- 6. Maintenance of confidentiality of records
- 7. Provision of free treatment for research related injury
- 8. Compensation of subjects for disability or death resulting from such injury
- 9. Freedom of individual to participate and to withdraw from research at any time without penalty or loss of benefits to which the subject would be entitled otherwise
- 10. Amount of blood sample (quantity in tea spoon full) to be taken
- 11. Costs and source of investigations, disposables, implants and drugs/ contrast media
- 12. Telephone number/ contact number of Principle investigator and Co-Investigator at the top of each page
- 13. In case of a drug trial:
 - a. The chemical name of the drug, date of its manufacturing and batch number must be mentioned
 - b. Initial bioequivalence study of the drug/references should be provided
- 14. Self-certification should be given that the translation to vernacular language is correct