

## 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, investigator must provide the subjects with the following information in English/ **Odia/ Hindi, in a simple layman's language, in a narrative form, directed to Participant /LAR, covering all the points given on the website**, which can be understood by them:

- i) Title of the Study/Project
- ii) Aims and methods of the research.
- iii) Expected duration of the subject participation.
- iv) The **benefits to be expected** from the research to the subject or to others.
- v) Any risk to the subject associated with the study.
- vi) Maintenance of **confidentiality** of records.
- vii) Provision of free treatment for research related injury.
- viii) **Compensation** of subjects for disability or death resulting from such injury.
- ix) Freedom of individual to participate and to **withdraw from research** at any time without penalty or loss of benefits to which the subject would otherwise be entitled.
- x) Amount of blood sample in quantity, in Tea Spoon Full, to be taken should be mentioned.
- xi) Costs and source of investigations, disposables, implants and drugs / contrast media must be mentioned.
- xii) **Telephone number/contact number of Principal Investigator and Co investigator at the top of each page.**
- xiii) In case of drug trials:
  - a) The **chemical name of the drug, date of its manufacturing and batch number** must be mentioned
  - b) Initial Bio equivalent study of the drug / references should be provided
- xiv) Self certification should be given that translation to vernacular is accurate.