


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
NURSING RESEARCH CODE OF ETHICS

- Voluntary written consent of the human subject will be obtained.
- In case of children and mentally handicapped subjects-voluntary written informed consent of the parents/guardians will be obtained.
- The probable risk involved in the project will be explained in full details to the subjects/parents/guardians.
- Subjects/parents/guardians will be at liberty to opt out of the project at any time
- .Researcher must inform subject about the study
- Research must be good for the community and society
- Researcher must try to avoid injury to research project
- Researcher must be qualified to conduct research
- Subjects or the researcher can stop the study if any problem arises.
- Researcher must protect the vulnerable group and other study participants from harmful effects of the experimental interventions
- Consider carefully and accurately the patient's rights and ethical concerns, in cases of using the patients for educational purposes.
- Medical services should not be affected if client /patient or their family do not wish to cooperate in education of the students.


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- To improve the healthcare services, the quality of nursing training courses, existing guidelines and standards must be continually reviewed and revised.
- Researcher, who are involved in research, must pass the specific training courses, have knowledge about the research regulations, and be familiar with national, general, and specific ethical guidelines and be pledged to these regulations.
- Researcher should not use their professional positions to convince the client / patient to participate in the research project.
- Refusal of participation in the research project by the patient or his/her family should not influence the delivery of the nursing interventions.
- Researchers are obliged to treat participant fairly and equitably before, during and after the research.
- Clinical researcher should make effort to enhance the expertise and clinical capacities of nursing and midwifery students.
- The researcher is responsible for ensuring confidentiality and privacy of the research participants and the data obtained from them.



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