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| **NUREB**GUIDE | **Participant Information Letters (PILs) /** **Informed Consent** Adapted from TCPS2, Chapter 3 |

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| **Purpose of Informed Consent** | To obtain assurance that participants in research projects clearly understand what they are agreeing to do, that they are free to decline involvement or withdraw from the study at any time, without penalty, and that all measurable steps are being taken to protect them. This process is meant to emphasize *Respect for Persons*. Consent **must** be obtained from participants **prior** to the recruitment of participants and conduct of research. |
| **Responsibility** | As per the Nipissing University Research Ethics Committee (NUREB) Policies and Procedures and the Tri-Council Policy Statement: Ethical Conduct for Research Involving Humans, researchers are responsible to ensure that the following information is included in your consent letter. |
| **Language** | Please use non-technical plain language. The letter should consistently be addressed to the reader (i.e., use “you”, “your”). |
| **Elements that must be included** | 1. Nipissing University logo
2. Names and contact information of investigator(s) and their department affiliation including daytime phone numbers and email addresses. For student research, the name and contact information of the Faculty Supervisor must also appear
3. The identity of the funder or sponsor, if applicable
4. Information that the participant is being invited to participate in this research project
5. The role of the research participants, including all procedures in which they will be involved (e.g., paper and pencil tasks, interviews, surveys, questionnaires, physical or physiological tests, etc.). Note if audio or other recording devices will be used
6. Length of time that will be required for each procedure or task
7. Frequency and overall duration, including the time associated with follow-up studies
8. The measures which you propose for providing feedback to research participants concerning the outcome of the research and any foreseeable secondary uses of the data (e.g., other studies, publications, etc.)
9. A plain-language description of all reasonably foreseeable risks (I.e., physical risks, discomforts or inconvenience as well as any psychological or social discomforts) that may be associated with participation in the research. If none, a statement that there are no known risks to participating in the study
10. Benefits to the research participants from their participation in the project
11. Benefits to society or to the advancement of knowledge from their participation
12. If incentives, remuneration, or compensation of any form will be offered to participants, then provide full details regardless of whether or not you are providing compensation
13. Describe the level of privacy, confidentiality or anonymity promised to participants
14. A statement to the effect that, by consenting, participants have not waived any rights to legal recourse in the event of research-related harm
15. **Right to withdraw** at any time without penalty or consequence; Participants are under no obligation to participate; are free to withdraw at any time without prejudice to pre-existing entitlements
16. **For focus groups**, a statement that confidentiality cannot be guaranteed if participants choose to speak outside the context of the research
17. **The Ethics Clearance statement**: This study has been reviewed and received ethics clearance through Nipissing University’s Research Ethics Board. If you have questions regarding your rights as a research participant, contact: Research Coordinator, Nipissing University, 100 College Drive, North Bay, ON P1B 8L7 or ethics@nipissingu.ca.
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| **Sample Statement:** Identified Participants | As a participant in this research project, I clearly understand what I agree to do. I am free to decline involvement or withdraw from this project at any time. I understand that steps are being taken to protect my safety and confidentiality of my data. I have read this Consent Form and have had any questions, concerns or complaints answered to my satisfaction. I confirm that I have been provided with a copy of this letter. |
| **Sample Statement:** Anonymous Participants | Any information that is obtained from you in connection with this study is anonymous. Participation in this study is voluntary. You are free to withdraw at any time. However, you may not be able to withdraw your data due to the anonymous nature of the participation. You have the right to refuse to answer any question(s) to which you object or that make you feel uncomfortable. Completion of this survey signifies your informed consent. Please keep a copy of this information letter for your records. |
| **Sample Statement:** Parental or Legal Guardianship Consent (for minors) | As a parent or legal guardian of the child participating in this research study, I clearly understand what I agree to do. I am free to decline my child’s involvement or withdraw them from this project at any time. I understand that steps are being taken to protect my child. I have read this Parent(s) or Legal Guardian(s) Consent Form and have had any questions, concerns or complaints answered to my satisfaction. I confirm that I have been provided with a copy of this letter.1. Right of the parent and participant to have their child’s personal information held confidential
2. Parental consent is not applicable
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| **If written consent is impossible** | Where written consent is impossible (Refer toTCPS2, Article 3.12) you **must**provide a written statement to the participants including all information that pertains to your research. Examples may include but are not limited to cultural appropriateness or sensitivities that may be perceived as an attempt to legalize or formalize the consent process, and therefore, may be interpreted by the participant as a lack of trust*.* |