## Instructions:

## Complete the assent form based on the research protocol.

## Some items in this template are optional or based on certain types of studies. Review the guidance throughout the template.

## Refer to HRP-013 Legally Authorized Representatives, Children, and Guardians, HRP-090 Informed Consent Process for Research, HRP-091 Written Documentation of Consent in the [HRPP Toolkit Library](http://research.umn.edu/irb/toolkit.html).

## For additional resources on drafting a consent form, consider the following:

## Medical terms, procedures, and conditions for [younger children](http://kidshealth.org/en/kids) and [teenagers](http://kidshealth.org/en/teens)

## [Plain Language Thesaurus for Health Communications](http://www.plainlanguage.gov/populartopics/health_literacy/Thesaurus_V-10.doc)

## [Clear Language and Design](http://clad.tccld.org/wp-content/uploads/2014/12/CLAD-Thesaurus.pdf)

## [MN Health Literacy](http://healthliteracymn.org/)

## [Readability calculator](http://www.online-utility.org/english/readability_test_and_improve.jsp) (use Flesch-Kincaid score)

## Remove help text and instructions before submitting the assent form draft to the IRB.

## The readability of the assent has to be age appropriate which means it could be at a 1st or 3rd or 6th grade reading level or anything in between depending on the age or circumstances of the minor. For example, detained adolescent youths typically have a reading level of 3rd grade. Do not assume that literacy and age are correlated.

## Young people are often confused by the “we” in the consent template. Unless there are a number of people involved in obtaining assent, use the pronoun ‘I,’ not ‘we.’

**University of Minnesota**

**Assent to Participate in Research**

## Title of Research Study: Study Title

## Researcher: Name of Investigator

**Sponsor:** Name of Sponsor

## What is research?

## Doctors and researchers are committed to your care and safety. There are important differences between research and treatment plans:

## The goal of research is to learn new things in order to help groups of kids in the future. Researchers learn things by asking a question, making a plan, and testing it. [For medical studies, otherwise delete.] In some kinds of research they are trying to find a new ways to help kids feel better. They do not know if the new medicine or treatment will work.

## The goal of treatment is to help you get better by using medication, therapy, surgery or other things that usually makes kids feel better. Sometimes treatments help make you feel better or get rid of the condition completely. Doctors can make changes to your treatment plan as needed.

## Why am I being asked to take part in this research study?

A research study is usually done to find a better way to treat people or to understand how things work. You are being asked to take part in this research study because you are [insert description of why they are being asked to take part].

## What should I know about being in a research study?

You do not have to be in this study if you do not want to do so. It is up to you if you want to participate and if you want to, talk to your parents about any questions or concerns you have about the study. You can choose not to take part now and change your mind later if you want. If you decide you do not want to be in this study, no one will be mad at you. You can ask all the questions you want before you decide.

If you become an adult (turn 18 years old or meet the legal definition of an adult found in Policy HRP-112) during this study, we will ask you if you want to continue to be in this study as an adult (Delete if not applicable).

## Why is this research being done?

In this study, I want to find out more about [insert purpose of the study in words understandable to the child].

## How long will the research last?

I expect that you will be in this research study for one session that will last [insert duration.]

## What happens if I say “Yes, I want to be in this research”?

If it is okay with you and you agree to join this study, you will be asked to [insert description or list of tasks they will be asked to complete (i.e. survey, tests, interviews, etc.).]

## Is there any way being in this study could be bad for me?

## [Delete this section if there are no risks or discomforts.]

## [The risks of procedures may be presented in a table form.]

## [Describe each of the following risks, if appropriate. If known, describe the probability and magnitude of the risk.]

## ● Physical risks

## ● Psychological risks

## ● Privacy risks

## ● Legal risks

## ● Social risks

## ● Economic risks

## ● Group or community risks

[Studies that include questionnaires or interview questions about mental health, psychological functioning, or mood, or includes participants that are at elevated risk of suicide, must include the following or similar language that is appropriate for the age and maturity of the child. Otherwise, delete.]

“As part of the research, we may ask questions about how you feel mentally and emotionally. We are providing a list of resources to you in case you would like to talk to someone and get help.  If you are thinking about hurting yourself or someone else, please tell someone who can help immediately. Call the toll-free 24-hour National Suicide Prevention Lifeline at 1-800-273-TALK (1-800-273-8255) to talk to a counselor near you.”

## Important information for girls [Remove if not applicable]

## [Include this information when appropriate for studies including girls that have reached puberty and there are risks related to pregnancy or breastfeeding. Otherwise remove.]

Because of the possible risk, you cannot participate in this study if you are pregnant or breastfeeding. You will have a pregnancy test before you begin any part of the study. If the test shows that you are pregnant, you cannot participate in this study.

Your parent/guardian will not be told the results of the pregnancy test without your permission. But, if your doctor believes that being pregnant may cause serious problems for your health, they may be forced to tell your parent/guardian the pregnancy test results.

If you are sexually active, you must agree to use an approved method of birth control during the study. Your study doctor or nurse can discuss acceptable methods of birth control with you.

You have the right to choose not to sign this form for any reason. If you do not sign, you cannot participate in this study.

## Important information for boys [Remove if not applicable]

## Include this information when appropriate for studies including boys that have reached reproductive potential. Otherwise remove.]

If you are sexually active, you must agree to use an approved method of birth control during the study. Your study doctor or nurse can discuss acceptable methods of birth control with you. Your parent/guardian will not be told.

You have the right to choose not to sign this form for any reason. If you do not sign, you cannot participate in this study.

## What happens to the information collected for the research?

The researchers will share your information, including research study records, to only people who have a need to review this information. For example, sometimes researchers need to share information with the University or other people that work in research to make sure the researchers are following the rules. The researchers may publish the results of this research or share the resulting data. However, we will keep your name and other identifying information confidential.

[Studies that include questionnaires or interview questions about mental health, psychological functioning, or mood, or includes participants that are at elevated risk of suicide, must include the following or similar language that is appropriate for the age and maturity of the child. Otherwise, delete.]

Anonymous Surveys (results are NOT individually identified):“We will not be able to link your responses to you, so we will not be able to provide you with personal feedback or referrals based on your responses to questions. If you are concerned about your mood, please refer to the attached resource referral information sheet. Please tell someone who can help right away. You can call also call the toll-free 24-hour National Suicide Prevention Lifeline at 1-800-273-TALK (1-800-273-8255) to talk to a counselor near you.”

Results Individually Identifiable/In-Person:“The study team may break confidentiality in an effort to keep you safe, which may include informing parents, local authorities, and/or health care professionals.”

“If you tell us that you are thinking about hurting yourself or others, the research staff may ask more questions. Depending on how intense your thoughts are or how much you feel like hurting yourself or others, the research staff may give you referrals for treatment, work with you to contact a personal doctor, trusted family member, or therapist to discuss your thoughts of harming yourself. We may need to work with you on a plan that might include getting to a medical facility for safety.”

## What else do I need to know?

If you agree to take part in this research study, the researcher will [describe any compensation that will be given to the child, when appropriate.]

## Who can I talk to?

For questions about research appointments, the research study, research results, or other concerns, call the study team at:

|  |  |
| --- | --- |
| Researcher Name:  Researcher Affiliation:  Phone Number:  Email Address: | Study Staff (if applicable):  Phone Number:  Email Address: |

To reach the research team: Please see the “Investigator Contact Information” section above.

To reach someone outside of the research team: This research has been reviewed and approved by an Institutional Review Board (IRB), a group of people that look at the research before it starts. This group is part of the Human Research Protection Program (HRPP). To share concerns privately with the HRPP about your research experience, call the Research Participants’ Advocate Line at [612-625-1650](tel:(612)%20625-1650) (Toll Free: 1-888-224-8636) or go to [z.umn.edu/participants](file:///C:\Users\cjarboe\Downloads\z.umn.edu\participants). You are encouraged to contact the HRPP if:

● Your questions, concerns, or complaints are not being answered by the research team.

● You are having difficulty reaching the research team.

● You want to talk to someone besides the research team or your parents.

● You have questions about your rights as a research participant.

● You want to get information or provide feedback about this research.

## Optional Elements:

## [Revise according to the study]

## The following research activities are optional, meaning that you do not have to agree to them in order to participate in the research study. Place your initials by each statement below to let us know your willingness to participate in these activities that may be required or optional.

|  |  |  |
| --- | --- | --- |
| **I agree** | **I disagree** |  |
| \_\_\_\_\_\_\_ | \_\_\_\_\_\_\_\_ | The researcher may audio or video record me to help do the research. The researcher will not share these recordings with anyone outside of the immediate study team, University, or other people that need to for the research. |
| \_\_\_\_\_\_\_ | \_\_\_\_\_\_\_\_ | The researcher may audio or video record me for use in scholarly presentations or publications, like a journal article. My identity may be shared as part of this activity, although the researcher will attempt to limit the ability to identify me. I understand what it means if my identity in some way is shared with others. |
| \_\_\_\_\_\_\_ | \_\_\_\_\_\_\_\_ | I agree to take a pregnancy test and/or use birth control if I am sexually active and agree to tell my doctor if I become pregnant. |

**Signature Block for Child Assent**

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Signature of child Date

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Printed name of child

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Printed name of person obtaining assent Date

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Signature of person obtaining assent