Consent to participate in a research study should be understood as a process rather than an event. The primary focus of ethical concern should always be on the quality of the consent process. As part of the informed consent process, the consent document is designed to provide information to potential subjects about a research study so they can make an informed decision about their participation. Documentation of the consent process is required unless specifically waived by the Institutional Review Board (IRB). One of the most common reasons for delay in IRB approval is incomplete, inaccurate, and/or unclear consent documents.

GENERAL REQUIREMENTS FOR INFORMED CONSENT [45 CFR §46.116]

Purpose of the Study

Describe the purpose(s) of the research study in lay terms. Include a statement that indicates why this is considered a research study. Provide definitions for specific research design features (e.g., randomization, longitudinal, causation).

Procedures

Provide a thorough description of the specific procedures involved in the study including which procedures are considered experimental and why. Include inclusion/exclusion criteria and length of involvement. If the subject will be interviewed or asked to complete a questionnaire, describe the types of questions that he/she will be asked to answer.

Compensation, Costs and Reimbursement

State whether compensation will be provided. If subjects will be compensated for their participation/reimbursed, describe in detail the type of payment, amount, and terms including circumstances where partial or pro-rated compensation would be provided. Specify any costs to the subject that may result from participation in the study that will not be reimbursed. (NB: Reimbursement is money given to research participants that reflects out-of-pocket expenses associated with research participation (e.g., childcare, transportation, parking, etc.); Compensation is money or items given to research participants that acknowledges the time and effort they have provided in participating in the research.)

Risks

Include information on any reasonably foreseeable risks or discomforts to the participants (physical, psychological/emotional, social, or legal). If applicable, include a statement that the treatment or procedure may involve risks, which are currently unforeseeable, to the subject (or to the embryo or fetus, if the subject is or may become pregnant).

Benefits

Describe all expected benefits and who will benefit. In some instances, it is acceptable to say that there are no direct benefits to subjects. Compensation for participation is not a benefit; provision for free drugs or procedures is not a benefit.
Withdrawal or Termination from Study

When applicable, subjects should be informed of circumstances under which their participation may be terminated by the investigator without the subject’s consent. Subjects should also be informed of procedures for safe and orderly termination should they decide to withdraw from the study before it is completed. Any conditions on withdrawal of data if participant chooses to withdraw from the study should be clarified (e.g., focus group discussions).

Confidentiality

Include information about the protection of subject’s privacy, method of protecting research data, and who may have access to study records. If relevant, different degrees of confidentiality should be described. Information regarding the use of audio and video recordings should be broken out as a separate option, to which participants can consent or not. Researchers should notify their subjects that in some instances, a representative of Office of Research Integrity may review research-related records for quality assurance in order to ensure that relevant laws and guidelines are followed. All information accessed by ORI will be held to the same level of confidentiality that has been stated by the research team. (NB: Confidentiality refers to the methods used by researchers to ensure that information obtained by them about their participants is properly protected. Privacy refers to how much control a person has over the circumstances of sharing oneself with others.)

Alternatives to Participation

Include applicable information on alternative procedures or courses of treatment that may be advantageous to the potential subject if he/she refuses to participate or withdraws from the study. Class-specific credit - if compensation for participation in the research is extra credit for a specific class, explain that the instructor is to provide a reasonable alternative way to earn extra credit. Psychology Department Subject Pool – if compensation for participation in the research is research credit, explain that an alternative to earning research credit is to attend a research lecture as described in the UCR introductory Psychology Research Participation Requirement.

Compensation for Injury

If applicable, standard non-alterable text describes the provision for subject injury incurred as a result of this study:

Social-Behavioral Research

If you are injured as a result of being in this study, the University of California will provide necessary medical treatment. The costs of the treatment may be billed to you or your insurer just like any other medical costs, or covered by the University of California or the study sponsor [sponsor name], depending on a number of factors. The University and the study sponsor do not normally provide any other form of compensation for injury. For further information about this, you may contact the UCR Office of Research Integrity via telephone at 951-827-5535 or via email irb@ucr.edu.

Biomedical-Clinical Research

If you are injured as a result of being in this study, the University of California will provide necessary medical treatment. The costs of the treatment may be billed to you or your
insurer just like any other medical costs, or covered by the University of California or the study sponsor [sponsor name], depending on a number of factors. The University and the study sponsor do not normally provide any other form of compensation for injury. For further information about this, you may contact the UCR Office of Research Integrity via telephone at 951-827-5535 or via email irb@ucr.edu.

Other Considerations

If a research team member has a disclosable financial interest in the outcome of this particular study or research program, a statement to that effect should be inserted. If the study involves collection of specimens, the appropriate standard language should be included. Some studies could generate findings that may affect participant's willingness to participate. Such findings should be disclosed to potential participants and this should be indicated in the consent form.

Contact Information

Include researchers’ contact information to answer study-related questions. Also, subjects should be instructed to contact the relevant UCR IRB if they have any concerns or questions regarding the study and/or their rights as research subjects: "If you have questions about your rights or complaints as a research subject, please contact the IRB Chairperson at (951) 827-4802 during business hours, or to contact them by email at irb@ucr.edu."

Voluntary Participation

Included a statement that emphasizes that the decision to participate, or not participate, is solely up to the subject. Subjects should be informed that they may decline or discontinue participation at any time without penalty or loss of benefits to which they are otherwise entitled.

Signature Lines

Signature and date line should be included for the subject. A “Legally-Authorized Representative” signature line should be included if you will obtain surrogate consent or are developing a parental permission form for enrollment of a minor. Also, you may need to obtain the assent for some minors in addition to parent’s consent.

Format and Style of Informed Consent Documents

- IRB requires that the consent and assent documents be written in the 2nd person, i.e., "you" rather than "I." Preferred language would be "The researchers are inviting you to be a part of a study..." Do not start sentences with "I understand..."

- Consent forms should be written in lay language, at a level understandable to the participants in the study. Researchers may use flowcharts and tables to enhance reading comprehension. Also, try to avoid medical/scientific/technical language or include simple explanations for such terms if they must be used.

- The use of a 12-point font is recommended. A larger type size may be appropriate for some populations, such as children, the elderly, or the visually impaired.

- Depending on the type of research and the methods of the study, differing terminology may be used to refer to those that are in the study: “subject,” “participant,” or even “student”
• The purpose of the research should be consistent with what was described in the IRB application. If the study is funded by a US federal entity, consistent with the funded grant.

• Consent document or script should have version dates to ensure that only the most recent and approved ones are being used.

• Researchers should stay away from using language that seems to indicate that the IRB has endorsed the research study.

• If applicable, a place for the subject to sign and date must appear on the consent document. Additional signature lines may be required if obtaining surrogate consent.

• The consent form should identify any external sponsor or funding agency, as well as the student’s and faculty supervisor’s affiliation.

• The informed consent, whether oral or written, may not include any language through which the subject or the representative is made to waive or appear to waive any of the subject’s legal rights, including release of the researcher, the sponsor, the university or its agents from liability for negligence.

• A summary of research results, and a mechanism to provide the summary, may be offered to participants.

Waiver of Written (Signed) Informed Consent [45 CFR §46.117(c)]

The IRB may waive the requirement to obtain a signed informed consent document in two situations:

• The only record linking the participant and the research would be the consent document;

• The principal risk would be potential harm resulting from a breach of confidentiality;

• Each participant will be asked whether the participant wants documentation linking the participant with the research, and the participant’s wishes will govern; AND

• The research is not a clinical investigation subject to FDA regulations.

OR

• The research presents no more than minimal risk of harm to participants, AND

• The research involves no procedures for which written consent is normally required outside of the research context.

In cases where the documentation requirement for informed consent is waived, the IRB often requires the researchers to provide participants with a written statement regarding the research. This written statement requires IRB approval.

Researchers interested in obtaining a waiver of written (signed) informed consent should make sure that their research qualifies for one of the above options, and should address how the research qualifies for each of the option’s requirements in their human subjects research application.
Waiver or Alteration of Informed Consent [45 CFR §46.116(d)]

The IRB may approve a consent procedure which does not include, or which alters, some or all of the elements of informed consent, or waive the requirements to obtain informed consent provided the researcher documents in their human subjects research application and the IRB finds that:

- The research involves no more than minimal risk to the participants;
- The waiver or alteration will not adversely affect the rights and welfare of the participants;
- The research could not practicably be carried out without the waiver or alteration;
- Whenever appropriate, the participants will be provided with additional pertinent information after participation.
- The research is not a clinical investigation subject to Food and Drug Administration regulations.

When an IRB waives the requirement to obtain informed consent, it waives the entire requirement for informed consent process. However, when the IRB grants an alteration of some or all of the elements of the informed consent (e.g., removes a required element of consent from the document), the process of obtaining informed consent is still required. Researchers interested in obtaining a waiver or an alteration of the consent process should select this as a procedure in their human subjects research application and address how the research qualifies for each of the above-listed requirements.