CONSENT TO PARTICIPATE IN RESEARCH

[Insert title of the study.] [If the study involves using different consent forms for different populations, identify the population group as the subtitle of the study.]

Suggested text:

You are asked to participate in a research study conducted at the [Insert the study site] by [name(s) of investigator(s)]. Your participation in this study is voluntary. You should read the information below, and ask questions about anything you do not understand, before deciding whether or not to participate.

Guidelines:

⇒ Use simple language.
⇒ Be concise.
⇒ Use the pronoun “you” consistently throughout (except for the signature of the subject on the last page).

• PURPOSE OF THE STUDY

[State what the study is designed to discover or establish.]

• PROCEDURES

Suggested text:

If you volunteer to participate in this study, we would ask you to do the following things:

Guidelines:

⇒ Describe the procedures chronologically using lay language, short sentences and short paragraphs. The use of table or flow diagrams will help to organize this section and increase readability. Distinguish which procedures are experimental and which are standard clinical treatments. Include screening evaluations and a listing of inclusion/exclusion criteria. Define and explain medical and scientific terms in ordinary language (for example, describing the amount of blood to be drawn in terms of teaspoons or tablespoons).
⇒ Specify the subject's assignment to study groups, the number of subjects expected to be enrolled, length of time for participation in each procedure, the total length of time for participation, frequency of procedures, location of the procedures to be done, etc.
⇒ For research involving randomization of subjects into different arms of studies, specify the randomization procedures.
⇒ For research involving the use of placebo, describe “placebo” in lay terms.
⇒ If pregnant women will be excluded, include HSPD Clause 009.01 or an equivalent statement on pregnancy prevention.

• POTENTIAL RISKS AND DISCOMFORTS

Guidelines:

⇒ Identify each procedure and then describe any reasonably foreseeable risks, discomforts, inconveniences, and how these will be minimized. Quantify risks using understandable comparisons.
⇒ In addition to physiological risks/discomforts, describe any psychological, social, or legal, risks that might result from participating in the research. If screening involves drug screening, serologic HIV or hepatitis C testing, explain the extent to which data will be kept confidential. Address local or federal reporting requirements, if any. Inform the subjects about availability of follow up or referral for treatment.

• ANTICIPATED BENEFITS TO SUBJECTS

Suggested text:

[Describe the anticipated direct benefits to subjects resulting from their participation in the research. If consent will be obtained from a legal representative of the subject, the direct benefit to the subject must be elaborated in the consent form. Refer to HSPD Clause 006.01, Title 10 United States Code, Section 980. ]

Guidelines:

⇒ If there is no likelihood that participants will benefit directly from their participation in the research, state as much in clear terms. For example: “You should not expect your condition to improve as a result of participating in this research” or “This study is not being done to improve your condition or health. You have the right to refuse to participate in this study.”
⇒ Do not include payment for participating in this section.

• ALTERNATIVES TO PARTICIPATION

Guidelines:

Date of preparation of current version:
Date of Approval:
Expiration Date:
⇒ Describe any appropriate alternative therapeutic, diagnostic, or preventive procedures that should be considered before the subjects decide whether or not to participate in the study. If there are no efficacious alternatives, state that an alternative is not to participate in the study.

⇒ If the prospective subjects are suffering from a terminal illness, and there are no alternative treatments available, you should say so; but add that treatment of symptoms and pain control are available through supportive care. In other words, avoid suggesting that participation in the research is the only way to obtain medical care and attention.

⇒ If prospective subjects have a chronic, progressive disorder, for which no treatment had been demonstrated to be safe and effective, say that, as well. But also describe opportunities for managing symptoms, improving ability to function, etc. so that it does not appear that the patient will be abandoned if he/she does not agree to participate.

• PAYMENT FOR PARTICIPATION

Guidelines:
⇒ Under 24 CFR 30, payment for participation to active duty military personnel is limited to blood donation and may not exceed $50 per blood draw. Active duty research subjects may not receive any other payment for participation in a research study.
⇒ State whether the subject will be paid or offered other benefits (e.g., free care). If not, state so.
⇒ If the subject will receive payment, describe remuneration amount, when payment is scheduled, and prorated payment schedule should the subject decide to withdraw or be withdrawn by the investigator.
⇒ If the subject will be reimbursed for expenses such as parking, bus/taxi, etc., state so.

• POSSIBLE COMMERCIAL PRODUCTS

(Note: If this does not apply to your research, please omit this entry and delete the heading.)

Suggested text:

HSPD Clause 004.01, Sample Donation.

Guideline:
⇒ If any human materials (tumor tissue, bone marrow, blood, etc.) will be stored for other uses or if samples are used for establishing a cell line which may be shared with other researchers and which may in the future be of commercial value,
subject must be informed of the fact in the consent form. and asked to sign a separate sample donation form.

- **MEDICAL CARE FOR RESEARCH RELATED INJURY**

Suggested text:

Note: The following is a required element of informed consent for research involving greater than minimal risk.

HSPD Clause 003.01, Medical Care for Research Related Injury.

- **CONFIDENTIALITY**

Suggested text:

When the results of the research are published or discussed in conferences, no information will be included that would reveal your identity. If photographs, videos, or audio-tape recordings of you will be used for educational purposes, your identity will be protected or disguised. [Describe how personal identities will be shielded, disguised, etc.]

[When the research records may be subject to inspection by FDA, a funding agency, or an industrial sponsor, you must add:]

Authorized representatives of the U.S. Army Medical Research and Materiel Command, FDA, and the manufacturer of the drug [or device] being tested [insert name of company] may need to review records of individual subjects. As a result, they may see your name; but they are bound by rules of confidentiality not to reveal your identity to others.

Guidelines:
⇒ Give a brief description of how personal information, research data, and related records will be coded, stored, etc. to prevent access by unauthorized personnel.
⇒ Explain how specific consent will be solicited, if any other uses are contemplated.
⇒ If applicable, state if and when individual responses to survey questionnaires will be destroyed, following analyses of the data.

- **PARTICIPATION AND WITHDRAWAL**

Suggested text:

Your participation in this research is voluntary. If you choose not to participate, that will not affect your relationship with (enter study site) or your right to health care or other
services to which you are otherwise entitled. If you decide to participate, you are free to withdraw your consent and discontinue participation at any time without prejudice.

- **CONSEQUENCES OF WITHDRAWAL**

(Note: If this does not apply to your research, please omit this entry and delete the heading.) [Explain the consequences of a subject's decision to withdraw from the research and any follow up the subject may be asked to complete, for reasons of safety.]

- **WITHDRAWAL OF PARTICIPATION BY THE INVESTIGATOR**

Suggested text:

The investigator may withdraw you from participating in this research if circumstances arise which warrant doing so. If you experience any of the following side effects [list and describe] or if you become ill during the research, you may have to drop out, even if you would like to continue. The investigator will make the decision and let you know if it is not possible for you to continue. The decision may be made either to protect your health and safety, or because it is part of the research plan that people who develop certain conditions may not continue to participate.

- **NEW FINDINGS**

During the course of the study, you will be informed of any significant new findings (either good or bad), such as changes in the risks or benefits resulting from participation in the research or new alternatives to participation, that might cause you to change your mind about continuing in the study. If new information is provided to you, your consent to continue participating in this study will be re-obtained.

- **IDENTIFICATION OF INVESTIGATORS**

In the event of a research related injury or if you experience an adverse reaction, please immediately contact one of the investigators listed below. If you have any questions about the research, please feel free to contact [Identify the point of contact. Include the daytime telephone numbers and addresses.]

For greater than minimal risk studies, include night/emergency telephone numbers.]

- **VOLUNTEER REGISTRY DATA BASE REQUIREMENTS**

For all studies involving greater than minimal risk to subjects, HSPD Clause 002.01, Volunteer Registry Data Base must be included in the consent form.

- **RIGHTS OF RESEARCH SUBJECTS**

Date of preparation of current version:
Date of Approval:
Expiration Date:
You may withdraw your consent at any time and discontinue participation without penalty. You are not waiving any legal claims, rights or remedies because of your participation in this research study. If you have questions regarding your rights as a research subject, you may contact the [Insert name, address, and telephone number of the point of contact.]

**SIGNATURE OF RESEARCH SUBJECT**
I have read the information provided above. I have been given an opportunity to ask questions and all of my questions have been answered to my satisfaction. I have been given a copy of this form.

________________________________________
Name of Subject

____________________________  ______________
Signature of Subject               Date

Address

**SIGNATURE OF WITNESS**
My signature as witness certifies that the subject signed this consent form in my presence as his/her voluntary act and deed.

________________________________________
Name of Witness

____________________________  _____________________________
Signature of Witness               Date (same as subject’s)