**Requirements for Ensuring Compliance with the Short Form Consent Process**

**For Non-English Speakers**

If your study targets a particular non-English speaking population, or if you expect to enroll more than 2 people of a specific non-English speaking population, you may be required to translate consent documents into that particular language. Please review the [IRB Policies and Procedures](http://www.irb.emory.edu/documents/PoliciesAndProcedures.pdf) for information regarding the translation policy.

Use of a short form is allowed when:

1. The Study Population page in eIRB includes “Subjects who are not able to clearly understand English”;
2. An Emory-provided short form is used or the IRB has approved a research team-provided short form;
3. Use is not expressly prohibited by the IRB; and
4. The study sponsor allows use of a short form.

If any of the above conditions are not met, an amendment requesting permission must be submitted and approved by the IRB prior to using a short form.

**Procedures for Using a Short Form:**

* No more than 2 short forms of the same language should be used for enrollment in a 12 month period. Any additional uses require consultation with the Emory IRB office.
* The **person obtaining consent** should ensure that contact information is noted on the short form in the blanks provided, with a name on the first line and phone number on the second line.
* A **translator** must read the English consent form and verbally translate the information to the subject or the subject’s legally authorized representative (LAR). If the subject is a child six years or older, the approved assent documents should also be verbally translated. The consent process must be witnessed by someone who is fluent in both English and the subject’s language. The **translator** may serve as the **witness** unless he or she is a member of the study team.
* A **witness,** who may also be the **translator** but cannot be affiliated with the study, must sign both the short form consent and the English consent (signing anywhere on the English consent signature page is acceptable).
	+ Studies with optional consent items: The **translator** must write a comment on the last page of the short form to indicate that the **subject** made specific choices. The **translator** should indicate the **subject’s** choices on the English consent form and include the **translator’s** initials beside each choice.
* The study **subject** or LAR must sign the short form consent (not the English version). If an LAR provides consent, it should be recorded as a note in the **subject’s** research record. If enrolling a child, the assent form is verbally translated but the child does not sign any documents.
* The **person obtaining consent** must sign the English version of the IRB-approved consent form.
* The study **subject,** or LAR, must receive copies of the following:
	+ The short form consent signed by the subject and the witness
	+ The IRB-approved English consent signed by the witness and person obtaining consent
* The original signed and dated IRB-approved English consent form should be filed *with* the original signed and dated short form consent in the **subject’s** research record.

**FOR REFERENCE ONLY – ENGLISH SPEAKING SUBJECTS MUST SIGN FULL IRB-APPROVED ENGLISH CONSENT FORM**

**Emory University**

**Consent to Participate in Research**

You are being asked to participate in a research study.

Before you agree, the investigator must tell you about (i) the purposes, procedures, and duration of the research; (ii) any procedures which are experimental; (iii) any reasonably foreseeable risks, discomforts, and benefits of the research; (iv) any potentially beneficial alternative procedures or treatments; and (v) how confidentiality will be maintained.

Where applicable, the investigator must also tell you about (i) any available compensation or medical treatment if injury occurs; (ii) the possibility of unforeseeable risks; (iii) circumstances when the investigator may halt your participation; (iv) any added costs to you; (v) what happens if you decide to stop participating; (vi) when you will be told about new findings which may affect your willingness to participate; and (vii) how many people will be in the study.

If you agree to participate, you must be given a signed copy of this document and a written summary of the research.

You may contact \_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_ at \_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_ any time you have questions about the research.

You may contact Emory University IRB at 404-712-0720 if you have questions about your rights as a research subject or what to do if you are injured.

Your participation in this research is voluntary, and you will not be penalized or lose benefits if you refuse to participate or decide to stop.

Signing this document means that the research study, including the above information, has been described to you orally, and that you voluntarily agree to participate.

\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_ \_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_

Signature of Participant Date

\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_ \_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_

Signature of Witness Date