RESEARCH INVOLVING PREGNANT WOMEN & FETUSES

Definitions

**Subpart B:** This term refers to the regulations which apply to research involving pregnant women and fetuses as subjects. Subpart B is found in 45 CFR 46 (DHHS).

**Pregnancy:** Encompasses the period of time from implantation until delivery. A woman shall be assumed to be pregnant if she exhibits any of the pertinent presumptive signs of pregnancy, such as missed menses, until the results of a pregnancy test are negative or until delivery.

**Fetus:** The product of conception from implantation until delivery.

**Dead Fetus:** A fetus that exhibits neither heartbeat, spontaneous respiratory activity, spontaneous movement of voluntary muscles, nor pulsation of the umbilical cord.

Federal Research Regulations

Federal regulations require additional safeguards when approving research involving pregnant women and fetuses. These special protections are found in Subpart B. The IRB may approve research involving pregnant women or fetuses only if all conditions for research are met (see description of conditions below) provided that the research also meets the general criteria for approval.

Description

The IRB must consider the general criteria for IRB approval for all studies, including those that involve pregnant women and fetuses. In determining whether the general criteria for approval are met, the IRB may have additional considerations for the research that would generally not be of concern in a normal adult population.

**Risks**

Risks that may be considered minimal when dealing with normal adults may be riskier when applied to pregnant women and fetuses. Efforts should be made to minimize any potential harm.

There may also be unforeseeable risks to an embryo or fetus for a particular treatment or procedure. The possibility of unforeseeable risks must also be taken into consideration.

Additional Protections for the Inclusion of Pregnant Women and Fetuses in Research

The IRB must consider the degree of risk and discomfort involved in the research in relation to the direct benefits it offers to the pregnant woman or fetus in order to determine whether the study is approvable under the federal regulations. The standard of review is conducted consistently, regardless of full board or expedited review. The IRB may approve studies involving pregnant women and/or fetuses only if the research fits into either category 1 or 2 below and also meets all of the following considerations 1-6: 45 CFR 46.204

**Category 1:**

The research holds the prospect of direct benefit for the woman or the fetus. Any risk to the fetus is caused solely by interventions or procedures that hold out the prospect of direct benefit for the woman or the fetus.

**Category 2:**

- The research holds NO prospect of direct benefit to the woman or the fetus, and
- The risk to the fetus is not greater than minimal, and

Please contact the IRB Office at (801) 581-3655 or irb@hsc.utah.edu for additional guidance.
The purpose of the research is to develop important biomedical knowledge which cannot be obtained by other means.

Additional Considerations (1-6):

(1) Where scientifically appropriate, preclinical studies, including studies on pregnant animals, and clinical studies on non-pregnant women have been conducted and provide data for assessing potential risks to pregnant women and fetuses;

(2) The risks to pregnant women and fetuses have been minimized, while still allowing the objectives of the research to be achieved.

(3) The individual(s) providing consent will be fully informed regarding the reasonably foreseeable impact of the research on the fetus.

(4) The individuals conducting the research are prohibited from offering inducements, monetary or otherwise, to terminate a pregnancy.

(5) The individuals conducting the research are prohibited from taking part in the decisions as to the timing, method, or procedures used to terminate a pregnancy.

(6) The individuals conducting the research are prohibited from determining the viability of a fetus.

Obtaining Consent for Research Involving Pregnant Women or Fetuses

Informed consent must be obtained from the necessary individuals as described below, unless the study meets the criteria to waive informed consent.

For research that falls into category 1 above, the board will make a determination based on the risk/benefit ratio as to which consent process method is appropriate for the study:

- If the research holds the prospect of direct benefit to the pregnant woman and fetus, consent must be obtained from the pregnant woman.
- If the research holds the prospect of direct benefit to the pregnant woman only, consent must be obtained from the pregnant woman.
- If the research holds the prospect of direct benefit to the fetus only, consent must be obtained from the pregnant woman and the father of the fetus.
  - The father’s consent does not need to be obtained if he is unable to consent because of unavailability, incompetence, temporary incapacity, or if the pregnancy resulted from rape or incest.

For research that falls into category 2 above, consent from the pregnant woman only is required.

Research on Dead Fetuses or Fetal Material (after delivery)

Research involving dead fetuses or fetal material obtained after delivery must be conducted in accord with any applicable federal, state, or local laws and regulations regarding such activities.

For more information about research involving the transplantation of fetal tissue, please see the Investigator Guidance Series: Transplantation of Fetal Tissue.

If information associated with the dead fetuses or fetal material is recorded in such a way that living individuals can be identified through the research, these living individuals are considered research participants and the researchers must abide by all regulations that apply to this group.

Additional Considerations

Please contact the IRB Office at (801) 581-3655 or irb@hsc.utah.edu for additional guidance.
VA Research:

Women known to be pregnant may be enrolled in research if all the requirements outlined in 45 CFR 46.204 are met including informed consent requirements and the following criteria:

1. Where scientifically appropriate, preclinical studies, including studies on non-pregnant women, have been conducted and provide data for assessing potential risks to pregnant women and fetuses;
2. The risk to the fetus is caused solely by interventions or procedures that hold out the prospect of direct benefit for the woman or fetus. If there is no such prospect of benefit, then the risk to the fetus is not greater than minimal and the purpose of the research is the development of important biomedical knowledge that cannot be obtained by any other means;
3. Any risk is the least possible for achieving the objectives of the research; and
4. The Facility Director certifies that the medical facility has sufficient expertise in women’s health to conduct the proposed research.

The VHA Office of Research and Development has additional guidance regarding collecting information about pregnancy progress and pregnancy outcomes for safety monitoring when pregnancy is not the research focus. More information may be found on the VA website (see link below).

Research in which the subject is a fetus, in-utero or ex-utero (including human fetal tissue) must not be conducted by VA investigators while on official duty, or at VA facilities, or at approved off-site facilities. Research related to in vitro fertilization must not be conducted by VA investigators while on official duty, or at VA facilities, or at approved off-site facilities.

Points to Address

New Study Application:

1. Participants page, question 3: Please select “Pregnant Women and Fetuses” as a vulnerable population. On the subsequent page (the Vulnerable Populations page) complete all questions justifying the inclusion of pregnant women and fetuses as participants in the study.

2. Risks and Benefits page: Describe all potential risks and benefits to pregnant women and fetuses.

Consent Document:

1. Describe the risks and benefits to the pregnant women and fetuses. State that there may also be unforeseeable risks to an embryo or fetus for a particular treatment or procedure.

References & Links

Additional Protections for Pregnant Women, Human Fetuses and Neonates Involved in Research (OHRP):

45 CFR 46, Subpart B

Investigator Guidance Series:

Transplantation of Fetal Tissue

http://irb.utah.edu/guidelines/investigator.php


Deleted: For research involving the participation of pregnant women as research subjects, the IRB must:↓

Determine that the proposed research meets the requirements outlined in this appendix;↓

Determine that adequate provision has been made to monitor the risks to the subject and the fetus.↓

Determine that adequate consideration has been given to the manner in which potential subjects are going to be selected, and that adequate provision has been made to monitor the actual informed consent process such as:↓

Overseeing the actual process by which individual consents required by this appendix are secured either by approving enrollment of each individual into the activity, or by verifying, perhaps through sampling, that approved procedures for enrollment of individuals into the activity are being followed, and↓

Monitoring the progress of the activity and intervening, as necessary, through such steps as visits to the activity site and continuing evaluation to determine if any unanticipated risks have arisen.↓

Research conducted at the VA which focuses on pregnant women may not be exempt.↓

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