



REQUIRED DOCUMENTS AND FORMS FOR IRB APPLICATIONS

The IRB reviews and approves supporting documents to the IRB application for the following purposes:

1. Information included in the document is necessary for the IRB to determine if the study meets the [Criteria for IRB Approval of Research](#), as well as other applicable federal regulations.
2. The document will be provided to the research participants and will assist them in making an informed choice about participation in the study.

Supporting documents that do not meet these purposes should not be submitted to the IRB for review and approval. Investigators are responsible for maintaining up-to-date versions of all supporting documents, including those reviewed and approved by the IRB and those that do not require IRB review.

This guidance document includes a list of supporting documents that are required to be included with the IRB application and meet the purposes described above.

New Study Applications

Attach required and supporting documents to the Documents and Attachments page

Social and Behavioral Science Projects	Biomedical and Health Sciences Projects
<ul style="list-style-type: none"><input type="checkbox"/> Method of Consent (and Authorization, as applicable) Documentation; may include one or all of the following:<ul style="list-style-type: none">o Consent and Authorization Documento Parental Permission and Authorization Documento Assent Documento Consent Cover Lettero Short Form and Written Summaryo Request for Waiver of Documentation of Consent and/or Authorization (form completed in the ERICA new study application, dependent on your response in ERICA)<input type="checkbox"/> Conflict of Interest Disclosure (completed electronically in the BRR system)<input type="checkbox"/> Recruitment materials, including advertisements or information intended to be seen or heard by potential participants<input type="checkbox"/> Full protocol (e.g. sponsor protocol), if the study is sponsor-initiated by industry, DHHS, etc.<input type="checkbox"/> DHHS-approved sample consent document (when one exists)<input type="checkbox"/> Grant application with budget but no appendices, for federal granting agencies	<ul style="list-style-type: none"><input type="checkbox"/> Method of Consent (and Authorization, as applicable) Documentation; may include one or all of the following:<ul style="list-style-type: none">o Consent and Authorization Documento Parental Permission and Authorization Documento Assent Documento Consent Cover Lettero Short Form and Written Summaryo Request for Waiver of Documentation of Consent and/or Authorization (form completed in the ERICA new study application, dependent on your response in ERICA)<input type="checkbox"/> Conflict of Interest Disclosure (completed electronically in the BRR system)<input type="checkbox"/> Recruitment materials, including advertisements or information intended to be seen or heard by potential participants<input type="checkbox"/> Full protocol (e.g. sponsor protocol), if the study is sponsor-initiated by industry, cooperative groups, DHHS, etc.<input type="checkbox"/> DHHS-approved sample consent document (when one exists)<input type="checkbox"/> For Investigational Drug and Biologic studies:

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



<ul style="list-style-type: none"><input type="checkbox"/> Questionnaires, interview questions and assessment instruments<input type="checkbox"/> Documentation of ancillary approvals from subcommittees for which the University of Utah has research oversight, such as:<ul style="list-style-type: none"><input type="checkbox"/> Protocol Review and Monitoring Committee (PRMC) – received electronically in ERICA<input type="checkbox"/> Conflict of Interest management plans (COI) – received electronically in ERICA<input type="checkbox"/> Institutional Biosafety Committee (IBC)<input type="checkbox"/> Primary Children's Hospital – received electronically in ERICA<input type="checkbox"/> Radiological Drug Research Committee (RDRC) – received electronically in ERICA<input type="checkbox"/> Resource for Genetic and Epidemiological Research (RGE) – received electronically in ERICA<input type="checkbox"/> Technology & Venture Commercialization Office (TVC)<input type="checkbox"/> Veteran Affairs Salt Lake City Health Care System (VASLCHCS)<input type="checkbox"/> Literature Cited/References<input type="checkbox"/> Principal Investigator's Scholarly Record (i.e. a curriculum vitae or resume)<input type="checkbox"/> Faculty Sponsor's Scholarly Record (i.e. a curriculum vitae or resume, if the principal investigator is not faculty)<input type="checkbox"/> Any other supporting materials relevant to the proposed research, if it exists	<ul style="list-style-type: none"><input type="checkbox"/> Investigator Brochure or Product Insert/Information Sheets<input type="checkbox"/> Investigational Drug Data Form (completed electronically in the ERICA new study application)<input type="checkbox"/> Documentation of IND receipt; may be documented by one of the following:<ul style="list-style-type: none"><input type="checkbox"/> FDA letter of IND receipt<input type="checkbox"/> A sponsor-generated document, such as the sponsor protocol, investigational brochure, or letter from the sponsor<input type="checkbox"/> For Investigational Medical Device studies:<ul style="list-style-type: none"><input type="checkbox"/> Verification of the IDE number; may be documented by one of the following:<ul style="list-style-type: none"><input type="checkbox"/> FDA letter providing the IDE<input type="checkbox"/> A sponsor-generated document, such as the sponsor protocol or letter from the sponsor<input type="checkbox"/> FDA letter granting an IDE for the proposed use or letter from sponsor stating that the study is a non-significant risk device study or letter explaining why the investigation is exempt from the IDE requirements under 21 CFR 812.2(c) or otherwise exempt<input type="checkbox"/> Grant application with budget but no appendices, for federal granting agencies<input type="checkbox"/> Questionnaires and assessment instruments<input type="checkbox"/> Documentation of ancillary approvals from subcommittees for which the University of Utah has research oversight, such as:<ul style="list-style-type: none"><input type="checkbox"/> Protocol Review and Monitoring Committee (PRMC) – received electronically in ERICA<input type="checkbox"/> Conflict of Interest management plans (COI) – received electronically in ERICA<input type="checkbox"/> Institutional Biosafety Committee (IBC)<input type="checkbox"/> Primary Children's Hospital – received electronically in ERICA<input type="checkbox"/> Radiological Drug Research Committee (RDRC) – received electronically in ERICA<input type="checkbox"/> Resource for Genetic and Epidemiological Research (RGE) – received electronically in ERICA<input type="checkbox"/> Technology & Venture Commercialization Office (TVC)<input type="checkbox"/> Veteran Affairs Salt Lake City Health Care System (VASLCHCS)<input type="checkbox"/> Literature Cited/References<input type="checkbox"/> Principal Investigator's Scholarly Record (i.e. a curriculum vitae or resume)<input type="checkbox"/> Faculty Sponsor's Scholarly Record (i.e. a curriculum vitae or resume, if the principal investigator is not faculty)<input type="checkbox"/> Any other supporting materials relevant to the proposed research, if it exists
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Please contact the IRB Office at (801) 581-3655 or irb@hsc.utah.edu for additional guidance.



Continuing Review Applications

Attach any updated/revised documents to the Update Study application, Documents and Attachments page, including new consent documents that need to be approved for another year.

- Currently approved Method of Consent (and Authorization, as applicable) Documentation; must include a clean copy for a new electronic stamp
 - If a study is permanently closed to new enrollment, it is not necessary to provide a currently approved consent form. The consent document will not receive a re-approval stamp (i.e., watermark) unless there is a specific request and explanation from the principal investigator or sponsor. This request must be approved by the IRB.
- Any other supporting material relevant to the proposed research, if it exists. This includes relevant multi-center trial reports, current risk-benefit assessments based on the study results, and participant benefits.

In addition, applications are required to submit the following, if applicable to the research:

- Documentation of ancillary approvals from subcommittees for which the University of Utah has research oversight, such as:
 - Protocol Review and Monitoring Committee (PRMC) – received electronically in ERICA
 - Conflict of Interest management plans (COI) – received electronically in ERICA
 - Institutional Biosafety Committee (IBC)
 - Primary Children's Hospital
 - Radiological Drug Research Committee (RDRC)
 - Resource for Genetic and Epidemiological Research (RGE) – received electronically in ERICA
 - Technology & Venture Commercialization Office (TVC)
 - Veteran Affairs Salt Lake City Health Care System (VASLCHCS)

Amendment Applications

Attach any updated/revised documents to the Update Study application, Documents and Attachments page.

- All previously-approved documents that are affected by the proposed changes must be submitted. A draft/tracked copy that highlights these changes as well as a clean copy are required for adequate review
- Any new documents created/required as a result of the proposed changes.
- Any other supporting materials relevant to the proposed changes, if it exists.

In addition, applications are required to submit the following, if applicable to the research:

- Documentation of ancillary approvals from subcommittees for which the University of Utah has research oversight, such as:
 - Protocol Review and Monitoring Committee (PRMC) – received electronically in ERICA
 - Conflict of Interest management plans (COI) – received electronically in ERICA
 - Institutional Biosafety Committee (IBC)
 - Primary Children's Hospital
 - Radiological Drug Research Committee (RDRC)
 - Resource for Genetic and Epidemiological Research (RGE) – received electronically in ERICA
 - Technology & Venture Commercialization Office (TVC)
 - Veteran Affairs Salt Lake City Health Care System (VASLCHCS)