Guidance and Procedure: IRB Review Level – Expedited Review
(last updated February 26, 2019)

**Introduction**

This guidance outlines the criteria and process for IRB review conducted by expedited procedures.

The IRB may review applications using Expedited review procedures if they meet specified criteria within the federal regulations, as noted below. An Expedited review may be performed by the IRB Chair and/or by an experienced IRB member designated by the IRB Chair, based on the member's area of expertise. In reviewing the research, the reviewer may exercise all of the authorities of the full Committee except to issue disapproval. The reviewer may at any time refer the application to the full Committee if necessary. All Expedited protocols are reviewed at least once annually.

**Categories of Research That May Be Reviewed By Expedited Review**

Protocols may be reviewed via an Expedited review process if they meet the following criteria, as listed in 45 CFR 46.110(b)(1):

- Research poses *no more than minimal risk to subjects*, as assessed by the reviewer; and
- Research for which *all of the procedures fall within the list of categories of research published by the Secretary of HHS* and the Food and Drug Administration (FDA).

Click here for November 9, 1998 Federal Register list of research eligible for expedited IRB review.

**General Restrictions:** Expedited review procedures may not be used where:

- Identification of the subjects and/or their responses would easily place them at risk of criminal or civil liability or be damaging to the subjects’ reputation, financial standing, employability, etc., unless reasonable and sufficient protections will be implemented so that risks related to invasion of privacy and/or breach of confidentiality are no greater than minimal.
- The Expedited review procedure may not be used for classified research involving human subjects.
Initial Review Process and Communications

Submission: Investigators are required to submit a complete application via webIRB, including all required supporting documentation.

Materials for review: All documents outlined in Materials Required for IRB Review are provided to designated expedited reviewers via webIRB.

IRB Pre-Review Process:

- IRB staff will conduct an initial pre-review for completeness, and determine whether a submission may qualify for Expedited review.
- The staff forwards the submission to the IRB Chair and/or designee for review.
- The reviewer makes the final determination of whether initial, continuing review, and modification submissions meet the eligibility criteria and falls into one or more of the categories listed above.
- The reviewer provides comments in written form.
- The IRB Chair/designee may request additional review by other member(s) of the IRB with applicable expertise. The additional assigned reviewer provides comments in written form.
- The assigned reviewer(s) of an Expedited submission may exercise all of the authorities of the IRB except that the reviewer(s) may not disapprove the research. A reviewer may choose to consult with another member prior to making any determinations. If the reviewer finds that the research should not be approved, it must be referred to the full Committee for final determinations.

Possible IRB Actions: The assigned reviewer(s) may make one of the determinations listed below:

- **Approved**: Acceptable as is. No changes are required. Criteria for IRB approval met.
- **Accepted Pending Modifications**: Minor specific changes are required.
- **Referred for Full Committee Review**: The reviewer may determine that the submission may not be expedited. The reviewer may also decide that additional information must be provided by the investigator prior to review by the convened Board.
- **Not Human Subjects Research**
- **Not Engaged in Human Subjects Research**

**NOTE**: Only the Full Committee may disapprove a study if the criteria for IRB approval are not met.

Communication with Investigators Regarding IRB Actions: Approvals, concerns and suggestions are communicated to investigators following each step of review; see IRB Review Process and Communication of Results of IRB Review for details.

Review Frequency: If the IRB Chair/designee determines that a protocol previously reviewed under Expedited review procedures requires review more often than annually, review of the protocol will be referred for review by the convened IRB. The IRB will communicate to the investigator in writing and document on the Approval Letter all determinations of requirements for review more often than annually.

Timely Review: The IRB and the staff aim to respond to the investigator within one week of receipt of the IRB submission. Depending upon the type of review (i.e., initial, amendment, continuation) expedited review approval may take from one to three weeks.

Documentation of Expedited Review Procedures: Regulatory determinations for initial and continuing reviews conducted under Expedited review procedures are documented in the webIRB application and on the IRB approval notices.
For protocols reviewed using expedited review procedures, the designated reviewer(s) designate the applicable expedited review category (ies) in webIRB when they electronically submit their Expedited Review Activity Form.

Expedited reviewers document any disagreement with the protocol-specific information supporting any waiver of informed consent or documentation of consent or the inclusion of vulnerable subjects in the research in the “Additional Comments” section of the Expedited Review Activity Form.

IRB staff include conditions on IRB approval notices to document the applicable expedited review category(ies), approved waiver(s) of informed consent or documentation of consent, and criteria for inclusion of vulnerable subjects in the research.

**IRB Approval Notice:** In addition, the Approval Notice will contain all regulatory determinations, the IRB approval period, and the list of funding sources for the research in addition to PI and study identifying information and other information as needed.

### Post-Initial Review

**Continuing Review:**

- **Pre-2018 Rule:** For federally-supported research reviewed and approved by Expedited Review procedures on or prior to January 20, 2019, the UCLA IRB must review ongoing research at least annually, unless the IRB requires more frequent review for a particular protocol.

- **Extended approvals issued prior to January 21, 2019:** The UCLA OHRPP/IRB issued approval for up to three years for non-exempt “research” involving “human subjects”, that involves no more than minimal risk to participants, and is not subject to federal oversight.

- **2018 Revised Common Rule:** For research reviewed and approved by Expedited Review procedures on or after January 21, 2019, unless an Expedited Reviewer determines otherwise, continuing review of research will not be required for the following:
  - Research eligible for expedited review in accordance with § .110;
  - Research reviewed by the IRB in accordance with the limited IRB review described in § .104(d)(2)(iii), (d)(3)(i)(C), (d)(7), or (d)(8);
  - Research that has progressed to the point that it involves only one or both of the following, which are part of the IRB-approved study:
    - Data analysis, including analysis of identifiable private information or identifiable biospecimens, or
    - Accessing follow-up clinical data from procedures that subjects would undergo as part of clinical care.

The UCLA IRB may elect on a study-specific basis, to apply the 2018 Revised Common Rule to studies that were approved prior to January 21, 2019.

For more information, please see [IRB Review Process – Continuing Review](#).

**Amendments to Approved Research:** All amendments to currently approved research must be approved by the IRB prior to implementation, except when necessary to eliminate apparent immediate hazards to the human subjects. For more information, please see [IRB Review—Amendments](#).
**Post-Approval Reporting:** The Principal Investigator must submit to the IRB all post approval reports that meet the submission criteria within the proper timeframe, as described in the OHRPP document [Post Approval Reporting Requirements (PAR)].

**Designation of Reviewers Other Than an IRB Chair**

- By virtue of the qualifications and experience necessary for the position, IRB Chairs and Vice Chairs are eligible to review on an Expedited basis. If needed to address considerations such as expertise, scheduling or submission volume, an IRB Chair or Vice Chair may identify other experienced members with sufficient experience and expertise to review on an Expedited basis. “Experienced” members must have active IRB service of at least one year and demonstrated knowledge and application of regulatory requirements and OHRPP/IRB policies and procedures.

- Designation of a member other than a Chair or Vice Chair as eligible to review according to the criteria for Expedited review procedures shall be confirmed via written communication among the following individuals as appropriate: the Chair, the designated member, the OHRPP Director or Assistant Director and the IRB Administrator. Any changes to the designation shall be communicated in the same fashion. The IRB Administrator shall maintain all documentation related to such designations. The IRB Administrators are responsible for communication with eligible reviewers regarding review duties. Because of their experience and qualifications, the IRB Assistant Directors and the IRB Administrators may serve as voting members and may be designated by the Chair to review Expedited studies as needed.

**References**

**DHHS Regulations**
- Expedited review procedures for certain kinds of research involving no more than minimal risk, and for minor changes in approved research: [45 CFR 46.110](#)
- Criteria for IRB Approval of Research: [45 CFR 46.111](#)
- November 9, 1998 Federal Register list of research eligible for expedited IRB review

**FDA Regulations & Guidance**
- Criteria for IRB Approval of Research: [21 CFR 56.111](#)
- Expedited review procedures for certain kinds of research involving no more than minimal risk, and for minor changes in approved research: [21 CFR 56.110](#)

**OHRP Guidance**
- [Guidance on Written IRB Procedures](#) – May 2018
- Desk Manual: Guidance for UCLA OHRPP Staff Designated to Conduct Review by the Expedited Procedures

**UCLA OHRPP Guidance**
- [Post Approval Reporting Requirements (PAR)](#)
- [IRB Review Process and Communication of Results of IRB Review](#)
- [Materials Required for IRB Review](#)

**Change history:**
- 2/26/2019: Updated to reflect 2018 Revised Common Rule requirements; updated OHRP link.