Part 3 – HRPP Policy

This section describes the process by which the University’s Human Research Protection Program (HRPP) policies, including the IRBMED policies, are developed, approved, and implemented, and articulates minimum requirements for IRBMED SOPs and Policies.

I. Rule making at the University of Michigan is divided three ways: (i) the Bylaws of the Board of Regents; (ii) rules initiated by University authorities that become effective only upon approval by the Board of Regents (Regents Policies); and (iii) rules adopted by subordinate University authorities, under delegated legislative powers, that become effective as provided by such subordinate authorities.

Human Research Protection Program (HRPP) policies fall within the third class of rule making. In Standard Practice Guide 303.05, the University has delegated to the Vice President for Research (VPR) general executive responsibility for the research programs of the University and, in that role, the responsibility for implementing the HRPP, including the legislative powers to adopt and enforce HRPP policy and procedures. See, SPG 303.05.

II. HRPP OPERATIONS MANUAL

The HRPP Operations Manual (OM) is the primary location for compiling, organizing, integrating, and pointing to the rules, policies, practices, and guidance encompassing the University’s HRPP. The VPR has approved the OM and approves each substantial modification or amendment to it. Records of such approval are maintained in the UM Office of Research (UMOR).

At least once every five years, typically in conjunction with the AAHRPP re-accreditation cycle, UMOR initiates a comprehensive review of the OM. Revisions may be made at any time, however, as required by changes in law, ethical standards, institutional policy, quality assurance activities, or other considerations. Non-substantive revisions (e.g., to correct typographical errors, update links, or incorporate summaries of new or revised laws or regulations governing the HRPP) may be made upon approval of the HRPP Director with notice to the VPR.

III. IRB STANDARD OPERATING POLICIES AND PROCEDURES

Refer to HRPP OM Part 3.II

A. General Provisions

The IRBMED members and staff to which these SOPs refer are accountable to the Medical School Associate Deans for Research and Regulatory Affairs and operate under the authority of UMOR with regard to the oversight of human subject research.

The IRBMED cooperates with the Medical School Associate Deans for Research and Regulatory Affairs, UM Health System (UMHS) Compliance Committee, and UMOR to establish, review, and revise these SOPs. These SOPs and any substantive revisions thereto, are subject to review and approval by the Medical School Associate Deans for Research and Regulatory Affairs, UMHS Compliance Committee, and the VPR. Non-substantive revisions such as modifications to enhance regulatory flexibility and workflows, inclusion of standard forms, guidance documents, and similar information developed by the IRBMED in consultation with the Medical School Associate Deans for Research and Regulatory Affairs and UMOR do not require further review or approval. Outdated sections of these SOPs will be archived in such a way that changes and dates of approval may be followed.

The IRBMED, in conjunction with Medical School Associate Deans for Research and Regulatory Affairs, UM Health System (UMHS) Compliance Committee, and the OVPR
maintains guidance documents on a number of topics targeted to the IRBMED, IRBMED Staff and researchers on the IRBMED website. In many cases the guidance expands on the information contained within these SOPs and are therefore referenced where appropriate.

IRBMED primarily has oversight of human subject research conducted by the Medical School faculty and staff or in the Health System facilities. Under certain conditions, oversight of a research project can be moved between IRBMED and the Health Science-Behavioral Science IRB (IRB-HSBS) with approval by the Chair of the Board (referred to in the SOPs as the “Chair”) and guidance from UMOR.

The IRBMED conducts its business through multiple IRBs, each of which is a separately registered IRB with the Office of Human Research Protections (OHRP) for purposes of University policy and the Federal-Wide Assurance (FWA).

<table>
<thead>
<tr>
<th>Board No.</th>
<th>Registration No.</th>
<th>FWA No.</th>
<th>Content</th>
</tr>
</thead>
<tbody>
<tr>
<td>A1</td>
<td>00000244</td>
<td>00004969</td>
<td>Biomedical</td>
</tr>
<tr>
<td>A2</td>
<td>00001996</td>
<td>00004969</td>
<td>Biomedical</td>
</tr>
<tr>
<td>B1</td>
<td>00001999</td>
<td>00004969</td>
<td>Biomedical</td>
</tr>
<tr>
<td>B2</td>
<td>00001995</td>
<td>00004969</td>
<td>Biomedical</td>
</tr>
<tr>
<td>C1</td>
<td>00005467</td>
<td>00004969</td>
<td>Biomedical</td>
</tr>
</tbody>
</table>

The IRBMED also provides review of cooperative group-sponsored projects through an agreement with the National Cancer Institute - Central Institutional Review Board (NCI-CIRB).

Refer to IRBMED SOP Part 5.II.A-D
Refer to HRPP OM Part 5.II.A-D

B. Organization and Personnel (Chairs, Members and Staff)

1. IRB Composition, Rosters, and Meeting Procedures

The IRBMED membership is selected to be sufficiently qualified through the experience, expertise, and diversity of its members (including consideration of race, gender, cultural background, and sensitivity to such issues as community attitudes), thereby to promote respect for its advice, counsel, and determinations in safeguarding the rights and welfare of human subjects.

Each of the five (5) registered IRBs consists of regular voting members and alternate voting members, with expertise augmented as necessary by consultants. New regular members first serve as alternate members for 6 month’s or other time period based on recommendation of the IRBMED Co-Chairs. An individual member may serve as a regular or alternate board member concurrently on more than one Review Board.

Each IRB will have at least five (5) voting members, including the Chairs, with varying backgrounds to promote comprehensive review of research activities commonly conducted at the Medical School and UM Health System (UMHS).

No IRB may consist entirely of members of one profession.

Every nondiscriminatory effort will be made to insure the IRB does not consist entirely of men or entirely of women. No selection will be made, however, solely on the basis of gender.

In addition to possessing the professional competence necessary to review specific research activities, each IRB that regularly reviews research involving one or more vulnerable categories of subjects, such as children, prisoners, pregnant women or physically or mentally disabled individuals will include members on the IRB of one or more individuals knowledgeable about and experienced in working with these subjects.
For projects supported by the U.S. Dept. of HHS’s National Institute on Disability, Independent Living, and Rehabilitation Research (NIDILRR) that “purposefully requires” inclusion of children with disabilities or individuals with mental disabilities, the IRB must include at least one member who is primarily concerned with the welfare of these research subjects (refer to 34 CFR 350 and 34 CFR 356).

When reviewing FDA-regulated studies, the IRB must include at least one physician.

The IRB must include at least one scientist member. A scientist is a member whose training, background, and occupation would incline him or her to view scientific activities from the standpoint of someone within a behavioral or biomedical research discipline. Scientist members include physician scientists (MDs or DOs), non-physician scientists (e.g., DDS, PhDs, nurses, geneticists, pharmacists and biomedical engineers), and social and behavioral scientists (e.g., psychologists, social workers, counselors). Scientist members have significant educational background (a science degree) and experience in scientific disciplines.

Any scientist who is an experienced regular member (i.e., those members designated as having enough experience to serve as Single IRB Member reviewers) or alternate member of an IRB may serve as a substitute chair of that Review Board in the absence of the appointed Co-Chair or Vice-Chair.

The IRB must include at least one non-scientist member. Non-scientist members are individuals without significant scientific educational background or experience and whose training, background and occupation would incline him or her to view scientific activities from a standpoint outside of any biomedical or behavioral scientific discipline. They may be recruited from active or emeritus University faculty or staff or from the community.

The IRB must include at least one member who is not otherwise affiliated with (or an immediate family member; spouse, domestic partner, or dependent) of a person affiliated with the University. The University supports efforts of the IRBs to include additional unaffiliated members.

“Unaffiliated” individuals include:

- University patients or research subjects or former students of the University who have no other affiliation with the University;
- Alumni, former faculty or staff of the University;
- Individuals contributing to fund-raising drives;
- Unaffiliated IRB members who have been paid at reasonable market rates for their services to an IRB.

“Affiliated” individuals include:

- Part-time employees;
- Current students;
- Members of any governing panel or board of the University;
- Paid and unpaid consultants of the University;
- Healthcare providers with medical staff membership or other credentials to practice at University clinical sites; and
- Volunteers working at the University on business unrelated to the University;
- Active emeritus faculty.
The IRBMED Staff maintains current membership rosters for each of the five (5) IRBs. Each membership roster contains a list of specified Chair(s), members and alternate members that are identified by name, earned degree, representative capacity (physician scientist, scientist, social-behavioral scientist, non-scientist); indications of experience sufficient to describe each member’s contributions to the IRB deliberations; and any employment or other affiliation or non-affiliation between each member and the University. Membership rosters are revised approximately quarterly to provide updates to include:

- new members or alternate members are approved by the Medical School Associate Dean for Regulatory Affairs;
- current members or alternate members extending their membership;
- members or alternate members who are moving from one IRB to another or serving on multiple IRBs;
- members or alternate members renewing membership after a period of time away from an IRB;
- members or alternate members that are resigning or are no longer eligible for membership.

The IRBMED will forward drafts of the revised membership rosters to the Medical School Associate Dean for Regulatory Affairs for review and approval. Following each roster change the Medical School approved membership rosters will be submitted to UMOR, which is then responsible for forwarding the approved membership rosters to OHRP. The IRBMED posts the current membership rosters of full and alternate on the IRBMED website.

a) IRB Chairs and Co-Chairs

Each Board has one Co-Chair and may have one or more Vice-Chairs (collectively referred to as the “Chairs” throughout these SOPs).

Each Chair must be a respected, active member of UM faculty, who qualifies as a scientist member with significant educational background, is concerned and knowledgeable about human rights and ethical issues, and is well informed concerning the laws, regulations, and University policies and procedures that govern the conduct of human subject research.

The Medical School Associate Dean for Regulatory Affairs is responsible for the appointment and reappointment of Chairs. When a vacancy arises, the Medical School Associate Dean for Regulatory Affairs may solicit nominations for a new Co-Chair or a Vice-Chair from the Medical School faculty, IRBMED members, staff, and consultants.

The Medical School Associate Dean for Regulatory Affairs then makes a recommendation to the UMHS Compliance Committee, which may adopt or reject the recommendation and in turn communicates its position to the Medical School Associate Dean for Regulatory Affairs.

An individual may serve an unlimited number of three (3) year terms as a Vice-Chair or a Co-Chair. In consideration of reappointment, a chair is evaluated for their contribution to leadership as well as their concern and knowledge of human rights and ethical issues, laws, regulations, and University policies and procedures that govern the conduct of human subject research.

b) IRB Members
Refer to HRPP OM Part 3.III.B.1

The Medical School Associate Dean for Regulatory Affairs may solicit nominations (including self-nominations) from members of the Medical School faculty, staff, and the University community. Unaffiliated representatives may be recruited by the Medical School Associate Dean for Regulatory Affairs or solicited through advertisements in local news media or by other means.

Solicitations may, as necessary, include information concerning the background, qualifications, and experience needed to promote diversity of experience and to provide or supplement necessary expertise on the IRBMED.

The Medical School Associate Dean for Regulatory Affairs will consult with the IRBMED Chairs on potential new members with regard to each individual’s qualifications, past participation (in the case of a reappointment), and other relevant criteria.

A potential new member will undergo an interview with the IRBMED Chairs and Medical School Associate Dean for Regulatory Affairs. The Medical School Associate Dean for Regulatory Affairs has final authority to make each member or chair appointment or reappointment.

All members should be sufficiently qualified through experience, expertise, and diversity and be able to determine the acceptability of proposed research in terms of institutional commitments and policies, applicable laws and regulations, and standards of professional conduct and practice.

Each member is appointed to an initial three-year term, which may be renewed at the discretion of the Medical School Associate Dean for Regulatory Affairs for an unlimited number of consecutive three (3) year terms.

Members are evaluated for reappointment at the discretion of the Medical School Associate Dean for Regulatory Affairs after seeking evaluation from the IRBMED Chairs and Office staff of the member’s level of participation, adequacy of reviews, regulatory/ethical interpretations.

Members will be assigned proposals to review that fall within their appropriate scientific and/or regulatory experience.

To ensure that the IRBMED is maintained as sufficiently diverse in experience, expertise, education, ethnicity, gender, cultural background, and sensitivity to such issues as community attitudes, the Associate Dean for Regulatory Affairs and IRBMED Chairs will periodically review the membership composition. Additional full or alternate members will be recruited to ensure sufficient breadth of the registered board composition should members’ terms expire, vacancies arise, or the submission review workload necessitate.

c) IRB Staff

The IRBMED is supported by a professional staff hired and supervised by the Director of the IRBMED. The Director reports to the Medical School Senior Associate Dean for Research through the Director of the Office of Research.

The Director and staff are responsible for facilitating IRBMED operations (human subject research application regulatory review; documentation and record retention; review of noncompliance allegations, including fact-finding; serving as an informational resource; conducting educational activities, etc.) in such a manner as to
maintain compliance with applicable State and Federal regulations and University policies, and for performing related activities as designated by the Medical School Senior Associate Dean for Research.

The Director assigns to each staff member the appropriate permission to perform regulatory and/or primary reviews; and/or coordinate the human participant research submissions in “eResearch”, i.e., the web-based eResearch Regulatory Management (eRRM) system which centralizes the review and approval process for Human Subjects Research Applications and IBC Biosafety Registrations.

The IRBMED Office includes a Regulatory Team for each of the five (5) IRBs. Each Regulatory Team is typically composed of a Senior Associate Regulatory Analyst (SARA), a Junior Associate Regulatory Analyst (JARA) and an Assistant Regulatory Analyst (ARA). The IRBMED office is further supported by other administrative, educational, compliance, quality assurance and quality improvement roles. All IRBMED staff ultimately report to the Director.

The IRBMED Office also tracks and manages membership information, including, but not limited to: membership role (physician scientist, non-physician scientist, non-scientist, and unaffiliated members), areas of expertise, COI, university affiliation, and advocacy for minority populations such as cognitively or physically disabled individuals, prisoners, and children or minors.

d) IRB Meetings

The Institution provides appropriate resources for board meetings including private meeting facilities equipped with appropriate electronic devices including overhead screens, projectors and individual notepad computers (issued for meeting use) for each member present that does not bring a personal laptop computer to the meeting. In addition, informational booklets containing copies of regulations are available at board meetings.

- The Co-Chair, or in his or her absence, a Vice-Chair or senior scientist member of the IRB leads each meeting. The IRBMED regulatory team monitors and documents attendance to ensure that the quorum, member composition, and diversity are present for each meeting as defined by Federal Regulations (21CFR56) A quorum (defined as more than half the number of regular voting members of an IRB must be present for each formal vote;
  - Quorum must include at least one non-scientist member;
  - At convened meetings at least one unaffiliated member who represents the general perspective of subjects should be present at the majority of meetings in a given year but is not required for quorum.
  - When reviewing research involving prisoners, the prisoner representative must be a voting member of the IRB.
  - If quorum is lost during a meeting, no voting will occur until quorum is restored.

e) IRBMED Meeting Schedules and Format

Each IRB convenes regularly to fulfill the mandate to oversee research involving human subjects subject to IRBMED’s jurisdiction. The IRBMED is comprised of five IRBs. IRBs A1, A2, B1 and B2 each convene twice monthly; two of the IRBs convene on weeks 1 and 3, and the other two IRBs convene on weeks 2 and 4. One Board (C1) convenes weekly to review studies with an oncology. Additional meetings may
be convened throughout the academic year, as necessary. The IRBs may meet by conference call only when necessary.

1) Meeting Cancellation

If circumstances dictate that a meeting should be cancelled (e.g., an anticipated lack of quorum), the IRBMED Regulatory Team will make a request of the Co-Chairs and the Director(s) to cancel the meeting after efforts to secure a meeting have failed. If the Co-Chairs agree the IRBMED office staff will notify the board members of the change. Agenda items will be reviewed for timely reassignment to other scheduled boards, if possible.

2) Ad Hoc Meetings

Occasionally additional board meetings are needed to address a significant increase in submissions or submissions from a previously cancelled meeting or other pressing issue. The Chairs are notified of the recommendation and asked for comment. If there is no disagreement, the members of the specific boards for which the additional meeting is necessary are notified by the IRBMED Regulatory Team to verify the availability of a quorum.

3) Alternate Board Meeting Format

In the event that a quorum of IRBMED members cannot be convened face to face, the IRBMED may utilize electronic technology (e.g., teleconference or videoconference) to facilitate the participation of the members. The agenda and all review materials will be available to the remote member via eResearch in advance of and throughout the meeting. The Chair of a meeting utilizing these alternative technologies will facilitate the active and equal participation of the remote members. Minutes of meetings utilizing assistive technology must document that these two additional conditions have been satisfied.

f) Agendas & Review Items

Prior to each convened IRB meeting the designated IRBMED Regulatory Team will prepare an IRB meeting agenda listing, and linking electronic items for review, discussion, deliberation; and vote, as appropriate; and other information of relevance to the IRB meeting’s purpose such as scheduled reports, i.e., Single IRB Member reviews (expedited, Single IRB Member for AE/ORIOs, or NCI-CIRB research projects). Updated working agendas are available at all times to IRBMED members, the Medical School Associate Dean for Regulatory Affairs, UMOR, and authorized consultants. The IRBMED Regulatory Team assigns incoming applications to upcoming meeting agendas generally based on the board assignment and availability of the designated primary reviewer.

Before a scheduled meeting, all IRBMED members are notified electronically of the planned meeting agenda generated by the IRBMED Regulatory Team. The agenda will contain links to all relevant items and documentation for review. IRB Members and Alternate Members will review the items attached to the agenda in advance of the IRB meeting. Primary reviewers will prepare a brief presentation of any submissions under their purview and recommendations for outcome.

IRB members and alternate members are encouraged to provide an RSVP to the IRBMED Office regarding their availability for each IRB meeting. Based on Member response the IRBMED Regulatory Team will review IRB meeting agenda items for
potential conflicts of interest with Members and provide recommendations for substitute Alternate Members.

g) Convened Meetings

(1) IRBMED Regulatory Team Responsibilities

On the day of, and in advance of each convened IRB meeting, the IRBMED Regulatory Team will oversee IRB meeting room facilities and supplies; the set-up of audio/visual projection equipment, laptop or notebook computers, and vote cards (as appropriate) for members, alternate members and others.

During the convened IRB meeting the IRBMED Regulatory Team will monitor attendance to ensure that quorum, member composition, and diversity are represented for each vote and record all voting outcomes and determinations on Voting Record sheets that are later archived in the IRBMED Office. The IRBMED Regulatory Team will provide support to the IRB by electronic projection of each agenda item for discussion, and any additional supporting or backup documents as needed.

The IRBMED Regulatory Team will also take meeting minutes to document IRB meeting attendance, reviews, discussion, outcomes, contingencies and determinations with regard to IRB meeting agenda items, federal and other state or local regulations, and IRBMED requirements.

Additional information will also be provided to all board members by the IRBMED Regulatory Team at the time of the convened meeting, including, but not limited to, continuing educational presentations and other relevant information to assist them in serving on the board.

(2) Changes to the Agenda

When an IRBMED meeting commences, all voting IRBMED members are alerted to any changes that may have been made to the planned agenda. In the event that an application is discussed that does not appear on the agenda (e.g., an emergency use or time-sensitive submission), a narrative summary of the protocol and sample informed consent forms, any recruiting materials, and other documents in the file are made available to all board members to review at the time of the meeting. Members will be afforded a reasonable period of time before a discussion and vote is taken.

(a) Board Member Reviews

The IRBMED Regulatory Team will assure that Board members have adequate time to review all aspects of the submissions for review. The twice monthly IRB meetings are scheduled for 3-5 hours which allows adequate time for assigned applications. The weekly (C1) board meets weekly for 2-3 hours which allows adequate time for assigned applications. However, if a board member feels they have been given inadequate time to review a specific submission then that item will be rescheduled to the next available meeting.

(b) IRBMED Deliberations and Voting

At the convened IRBMED meeting, the primary reviewer and any additional reviewer or consultant presents their review of the submission, including comments documented on their Review Checklist in advance of the meeting,
and their recommendation to the IRB, including any suggested changes. The IRBMED Regulatory Team and primary reviewer must complete their Reviewer Checklists prior to presentation to the convened IRB.

Following the primary reviewer’s presentation, board members discuss the submission and deliberate prior to voting.

An initial, amendment, or SCR submission may be approved or disapproved only upon a majority vote by the voting members present. The PI or study team designee may be requested to attend the convened board meeting in person or by telephone to address any questions raised by the board. However, neither the PI nor the study team will be permitted to be present for the discussion or vote of the submission.

h) IRBMED Meeting Minutes

(1) Content Requirements

Following a Review Board meeting, designated office staff shall prepare minutes consisting of at least the following information:

- Attendance of the members at the full convened board meeting, including a notation of absences of board members;
- Documentation of any conflicted members or staff;
- The time a full or alternate member leaves the room and rejoins the meeting;
- Acknowledgement of reviews approved by the Single IRB Member (expedited, Single IRB Member for AE/ORIOs, or NCI-CIRB research projects) review procedure;
- The names of PIs, guests and/or consultants in attendance;
- For each submission reviewed, any votes or other actions taken, and the vote on each action including:
  - the number and names of members voting for or against;
  - the number and names of those members abstaining;
  - the names of alternate members standing in the capacity of an absent regular member—based on designation: for example, non-physician scientist alternate member \( a \) is standing-in for non-physician scientist regular member \( b \);
- The names of conflicted members, consultant, PIs or guests who leave the room for the deliberation and vote.
- For initial and SCRs, the approval period;
- Protocol-specific information supporting any waiver of informed consent or documentation of consent, e.g. The waiver of HIPAA authorization or the inclusion of vulnerable subjects in the research;
  - Research involving pregnant women, fetuses and neonates;
  - Research involving prisoners; and
  - Research involving children
- The name of any consultant reviewer used for an application;
- The basis for requiring changes in or disapproving research;
- Separate deliberations for each action;
- A written summary of controverted issues and their resolution;
- A summary of any continuing education provided to IRB members;
- Documentation of expeditable studies that were reviewed by single-member reviewers prior to the meeting.
- Documentation of board deliberations and determinations involving
UaPs/UPIRSOs that must be made separately:
- An evaluation of unexpectness, in terms of nature, severity or frequency;
- An evaluation of relatedness;
- An evaluation of harm;
  - representative of potential increased risk to subjects or others; or
  - representative of risk of actual harm to subjects or others.
- When following DHHS regulations or guidance, documentation of justification of any deletion or substantive modification of information concerning risks or alternative procedures contained in a DHHS-approved sample consent document; and
- When following FDA regulations or guidance, documentation of the rationale for significant risk/non risk device determinations.

(2) Review and Ratification Process

Typically, within one (1) week (for the C1 board) but not more than four (4) weeks from the meeting date, an IRB’s DRAFT minutes shall be prepared. The DRAFT minutes are reviewed for quality, completeness and compliance with regulatory requirements by the IRBMED Regulatory staff and then sent electronically to Board members for review.

- Board members may request changes to the DRAFT if needed;
- IRB members will vote to ratify them at a subsequent meeting;
- The ratified minutes will be maintained in accordance with applicable legal requirements and institutional policy.
- In circumstances where the minutes require further scrutiny or review, i.e., after an outcome of serious and/or continuing noncompliance, the minutes may be presented to the board later than four (4) weeks after the meeting date. The Co-Chair of the board will notify the board members at the next available full convened board meeting should a delay in ratifying the IRB’s minutes is necessary.
- Minutes are archived in the eResearch System.
- In the event that minutes require amending due to a discovered error, for example, strict version control is applied to preserve the original minutes.

Additional guidance is available to IRB Regulatory Teams and staff regarding preparation, approval and amending IRBMED meeting minutes.

2. Use of IRB Consultants

The IRBMED, the Medical School Dean for Regulatory Affairs, or the VPR may, at their discretion, invite persons whose experience or expertise may aid the IRBMED in performing its responsibilities. Whether during meetings or otherwise, such individuals may include consultants, advisors, and ad hoc reviewers, or others who may serve the IRBMED, for example, by assisting in the review of a complex research project.

These individuals may participate in the discussions of, or provide written documentation concerning an application, but shall not be counted for the purposes of establishing quorum, nor shall they vote on the approval, disapproval, or other disposition of any application.
As appropriate, key information from consultants, advisors, and ad-hoc reviewers will be recorded in the minutes. These individuals will be granted access only to the assigned research project. Any individual asked to serve the IRBMED in this manner will be required to sign the standard IRBMED confidentiality agreement, follow the standard IRBMED member conflict of interest procedures, and comply with appropriate application review requirements.

Refer to [HRPP OM Part 3.III.B.2.](#)

### 3. Alternate IRB Members

The IRBMED may appoint alternate voting members to serve in the absence of regular voting members to establish quorum and participate in deliberations and votes on applications pending before the IRBMED. A regular voting member of one IRB (A1, A2, B1, B2, or C1) is automatically considered to be an alternate member to each of the other IRBs; specific designation on each of the rosters is not required.

Each alternate IRB member has experience, expertise, background, professional competence, and knowledge comparable to that of the primary IRB member whom the alternate would replace.

Alternate members may attend IRB meetings even when their attendance is not necessary to establish quorum. Alternate members may participate in the discussion; however, they may not vote unless designated to serve in the absence of a regular voting member.

A regular member from one IRB may serve as a primary reviewer on another IRB in the capacity of ad hoc reviewer and is not counted towards quorum or utilized as an alternate.

The IRBMED Chairs may reassign a voting member of one IRB as a voting member of another IRB, or may reclassify a regular voting member as an alternate voting member or vice versa, by notifying the member, the Medical School Associate Dean for Regulatory Affairs, and UMOR. When a reassignment of a voting member is made, i.e., a member moves from one IRB to another, the membership rosters are revised to reflect this change and the Medical School Associate Dean for Regulatory Affairs and UMOR are notified. The IRBMED Office maintains the membership information.

### 4. IRB Educational and Training Activities

Refer to [HRPP OM Part 13](#)

Refer to [IRBMED SOP Part 13](#)

#### a) Orientation of IRB Members

The IRBMED orientation program for new members is a multi-component program designed to permit new members to assimilate information in a manner that enhances retention and appropriate application of the material to reviews. The program includes workshops, directed mentoring, completion of human research educational modules in Program for Education and Evaluation in Responsible Research and Scholarship (PEERSS), the UMHS Health Insurance Portability and Accountability Act (HIPAA) training for researchers, and mock protocol reviews. Topics covered include but are not limited to:

- Human Subject Protections Overview
- Federal Regulations and the IRBMED Review Process
- Federal Regulations—Special Populations
- Review of eResearch Applications
New members are initially considered alternate members. At the conclusion of the orientation period a new member will be asked to conduct a mock review with the IRBMED Chairs and to present the review in the context of an IRB meeting. If the mock review is deemed to provide information in compliance with the regulations, the member enters a practicum period during which new members attend meetings and review protocols as primary reviewers but are not permitted to vote unless a regular member is for whom they qualify as an alternate is absent.

IRBMED Chairs determine when each new member’s cumulative experiences qualify them for appointment as a regular member and if they qualify to serve as a Single IRB Member reviewer (including expediting reviewer). This may occur at any time after presenting a mock review and usually occurs within one (1) year for bi-weekly members and six (6) months for weekly members.

b) Orientation of IRB Staff

Depending upon the role of the new staff member, completion of some or all of the IRBMED member orientation and continuing education workshops, as well as workshops offered to research personnel, are required at the discretion of the employee’s direct supervisor.

IRBMED staff members are required to complete all PEERRS human subject modules. Staff members are encouraged to attend local, regional, and national conferences on ethics, State and Federal laws, and regulations for human subject research protection.

c) Orientation of IRBMED Chairs

IRBMED Chairs are appointed per IRBMED SOP Part 3.III.B.1.a. Chairs meet with the Medical School Dean for Regulatory Affairs to review roles and responsibilities of the role in association with relevant federal and state regulations, laws, guidance materials, and University and Medical School policies. The Directors of IRBMED and Regulatory Affairs as well as relevant IRBMED Office staff also meet with the Chairs to provide orientation to the working procedures associated with the IRB. Feedback associated with the orientation sessions are provided to the Dean for Regulatory Affairs.

d) Continuing Education for IRB Members and Office Staff

All IRBMED Chairs, members, and staff participate in continuing education within the context of the IRB meeting and elsewhere. Continuing education on ethics, regulations, federal guidance, university policies, and eResearch are provided in the form of webinars, workshops and presentations at meetings. Printed and electronic materials are provided on an on-going basis.

U-MIC web-based tips are presented at each IRB meeting; annually, IRB Chairs, members and Office staff are offered an opportunity (the number of attendees varies based on budget) to attend national IRB meetings, IRB Seminar Series are presented to the research community, IRB Seminar Series are offered on specifically for board members on selected topics. IRBMED staff also participate in ongoing continuing education within the context of weekly staff meetings.

Reference materials such as U-MICs and the slides from Seminar Series are posted to the IRBMED website. IRBMED also prepares topic-specific guidance which is posted to the website.

e) Researcher Education
Refer to HRPP OM Part 13
Refer to IRBMED SOP Part 13

5. IRB Compensation and Liability Coverage

a) Compensation of Chairs

The IRBMED Chairs are compensated for the portion of their effort required to perform their duties as Chair. The IRBMED Chairs are paid a portion of their salary for the time and effort involved in performing the duties of a Co-Chair or Vice-Chair. The stipend is commensurate with the required time to perform the IRBMED duties, in negotiation with the Medical School Associate Dean for Regulatory Affairs or department Chair.

b) Compensation of Members of the Community

Unaffiliated members of the community are paid for their service on the IRB in an amount to compensate their time and expenses to attend the meeting, unless waived by the unaffiliated member. This may include, but is not limited to: parking expenses, computer support, and a per-meeting stipend. Members of the community that are deemed affiliated are offered compensation at the same rate.

c) Periodic Review of Compensation

The amount of compensation for Chairs and members of the community is reviewed periodically and may be changed by the Medical School Associate Dean for Regulatory Affairs.

d) Liability Coverage

Liability coverage is a matter of institutional policy and is further described in HRPP OM Part 3.III.B.5.

6. Evaluations of IRB Chairs, Members, Staff and Regular Consultants

Refer to HRPP OM Part 3.III.B.6.

a) Chairs and Member Performance Review

Annually, the Medical School Associate Dean for Regulatory Affairs will evaluate the Co-Chairs of the board to ensure that their expertise adequately addresses the types of protocols reviewed and to ensure that each Chair is an active participant and is trained in current interpretations of federal regulations and other relevant ethical principles for the protection of human subjects. The Vice-Chairs will be evaluated by the Co-Chairs at the conclusion of their first year of appointment as Vice-Chair. Thereafter, the Vice-Chair will be evaluated prior to renewal as Vice-Chair. A Vice-Chair may be evaluated more frequently if there is a concern as to their ability to fulfill the role. Feedback on the performance evaluation will be provided to the Co- or Vice-Chair along with any suggested corrective actions such as additional educational requirements or direction on how to improve workflows associated with the convened IRB meetings.

The IRBMED Co-Chairs and Vice-Chairs will evaluate new members of their boards at the conclusion of their first year of service to ensure that the expertise of each full and alternate member adequately addresses the types of protocols reviewed and to ensure that each member is an active participant who is trained in current interpretations of federal regulations and other relevant ethical principles for the
protection of human subjects. Feedback on the performance evaluation will be
provided to the member by the Co- or Vice-Chair along with any suggestions for
improving performance via additional education or mentoring.

Each board member reaching the end of their term is evaluated and provided
feedback by the Co-Chairs or Vice-Chairs prior to renewal. At any time during a
member's term an evaluation may be conducted if there is concern about the level of
a member's participation, adequacy of reviews, regulatory interpretations or other
ethical concerns.

b) Removal of a Chair or Member from an IRB

If necessary, the Medical School Associate Dean for Regulatory Affairs may relieve a
Chair or Vice-Chair from IRBMED service due to repeated non-attendance, lack of
participation in continuing education, or other problematic performance issues.
Should this action be required, the Medical School Associate Dean for Regulatory
Affairs will notify the Medical School Dean and UMOR.

Similarly, the Chairs may recommend to the Medical School Associate Dean for
Regulatory Affairs that a board member be relieved from IRBMED service due to
repeated non-attendance, lack of participation in continuing education, or other
problematic performance issues. In this situation, as described above for the Chair or
Vice-Chair, the Medical School Associate Dean for Regulatory Affairs will notify the
Medical School Dean and UMOR.

c) IRB Staff Performance Review

Staff members are evaluated yearly in a performance appraisal conducted by the
IRBMED Director and their functional supervisor as instructed by the IRBMED
Director. The IRBMED Director is evaluated by the Director of the Office of Research
in a yearly performance appraisal. If circumstances dictate, the Director and staff are
evaluated more often. Constructive feedback is provided to effectuate additional
learning or corrective action as necessary.

d) Regular Consultant Performance Review

In the rare event that an IRB regularly uses a consultant to conduct reviews, they will
be reviewed annually.

7. Conflicts of Interest involving Chairs, Members, Consultants and Staff

Refer to HRPP OM Part 9.II-III

It is the responsibility of IRBMED Chairs, members, consultants and staff to disclose both
actual and perceived conflicts of interest (COI) throughout their terms of service or
employment. The financial disclosure sections of the eResearch application indicate
disclosure of a financial interest in a sponsored project or technology transfer agreement.
This information is on file in the IRBMED office.

Disclosed COI information associated with IRBMED members and consultants is
obtained from UMOR and the Medical School’s COI Committee. This information is
considered during review assignment in order to ensure a member is not assigned to
review research for which they are identified as a conflicted member.

a) Financial Disclosures

At the beginning of their service and annually thereafter, each IRBMED member or
consultant completes an M-Inform disclosure for their financial disclosures. The
financial disclosure section divulges any significant financial interest in a sponsored project or technology transfer agreement. This information is periodically reviewed by the IRBMED office and relevant information is entered into eResearch. IRBMED members and consultants are also queried periodically as to any other relationships (e.g., familial) that might create a conflict of interest. This information is entered into eResearch.

Staff members in leadership or management roles, such as the Director, are required to complete an annual M-Inform disclosure or, more frequently as needed.

b) Conflicts of Interest with Research Involving Chairs and Members

An IRBMED member (including the Chair) is not assigned to review research if the member:

- Is a PI on the study or the PI's immediate relative (spouse, domestic partner or dependent);
- Has a significant financial interest in the research (as defined by University and Medical School policies on COI);
- Has other conflicts that the member, review board, Medical School Associate Dean for Regulatory Affairs, COI Committee, or UMOR believes might hamper the member's ability to perform an impartial review of the research.

Any conflicted reviewer (Chair or member) shall not be present for, count for quorum, participate in deliberations on, or vote on the disposition of research for which the individual has a conflict as described above. The conflicted reviewer or consultant may, however, be invited by the IRB to provide information relevant to the board's consideration of the research.

The conflicted reviewer or consultant must be absent from the room during both relevant deliberation and voting.

A conflicted Chair or member shall not participate in the investigation of actual or alleged noncompliance on behalf of the IRBMED (other than to cooperate with the investigation). if the conflict is as described above.

All conflicts of interest for studies reviewed at the convened IRB are documented by IRBMED staff in the IRB meeting minutes.

In some instances, an IRBMED member may have involvement in a research study that solely involves the provision of a service to a study (e.g., a Pharmacist from the Research Pharmacy who prepares and dispenses study medication, or a Radiologist who performs a diagnostic imaging study that is part of the research). The IRBMED does not consider this to be a conflict of interest with regard to reviewing an IRBMED submission, provided the member's role in the study is limited to providing a service to the PI and they are not otherwise engaged in the research study. For example, a board member is not permitted to be listed on the FDA Form-1572. If additional clarification is needed, contact IRBMED. This is consistent with the examples of non-engagement in research provided in OHRP's Guidance on Engagement of Institutions in Human Subject's Research, October 16, 2008, Section III.B.1.

c) Conflicts of Interest with Research Involving IRB Staff

An IRBMED staff member would be recognized as having a COI with research in which he/she has a significant personal or financial interest.

When a conflict is identified by UMOR, the University, the Medical School COI Board or other University unit (i.e., ORSP), and/or by self-disclosure, the IRBMED staff
person must excuse him/herself from administrative handling of the research and from the IRBMED board meeting where there is deliberation and vote on the research. IRBMED staff document all conflicts of interest in the IRB meeting minutes.

IRBMED staff shall not participate in the investigation of actual or alleged noncompliance or other misconduct if the staff person has a conflict as described above.

d) Conflicts of Interest Involving Consultants

Refer to HRPP OM Part 9.III.

Refer to IRBMED SOP Part 9.III

Refer to IRBMED SOP Part 3.III.B.7.b.

Consultants are not voting members of the IRB. No consultant to IRBMED may participate in the IRB’s review of an initial application, amendment or continuing review application, or participate in the investigation of actual or alleged noncompliance or other misconduct investigation in a research protocol, if a conflict of interest exists (as defined in Part 9 of the HRPP OM) except in limited circumstances as requested by the IRB. The consultant may be invited by the IRB to provide information relevant to the IRB’s consideration of the application taking into account the consultant’s special qualified expertise and their ability to provide an objective assessment. Any conflict will be disclosed to the board at the convened meeting prior to any participation of the consultant in the discussion.

The IRBMED Regulatory Team will evaluate whether an actual or perceived COI exists prior to contacting a consultant and also ask the consultant to disclose any perceived, potential, or actual conflicts. The Medical School COI Committee or UMOR will be consulted, if needed. Conflicts of interest involving consultants will be evaluated according to the same definition as applied to IRBMED members.

e) Conflicts of Interest Involving Guests

For guests attending a board meeting during the course of which a conflict is identified either by the IRBMED staff, board members, and/or by self-disclosure, the guest will leave the room during the discussion and vote on the research protocol. IRBMED staff will document the name of the guest, conflicted project(s), and the time the guest leaves and returns to the meeting.

f) Conflicts of Interest Involving a Convened Board

Prior to each convened IRBMED meeting, the IRBMED Regulatory Staff will determine if, to the extent possible, if a COI statement is documented for any research submissions for convened IRB review and will note the conflict on the agenda. However, it is ultimately the responsibility of the board member to self-identify any COI at the time it is known.

No IRBMED member, including the Chairs, shall be present for or participate in, the deliberations or vote on the disposition of an application for which the member has a conflict as described above. The member may, however, be invited by the IRBMED to provide information relevant to the board’s consideration of the application.

IRBMED Chair and staff will ensure that all identified, conflicted IRBMED members are:

• excused from discussion, except to provide information requested by the IRBMED; excused from deliberation;
• excused (absent from the room) during voting;
• not counted towards quorum for a particular vote; and
• documented appropriately in the meeting minutes

To facilitate the identification of any previously unreported conflicts, the IRBMED Chair shall, at each meeting, inquire whether any members should excuse themselves from discussion and voting as outlined above.

g) Conflicts of Interest Involving Single Member Review and Expedited Review

Prior to a Single Member review (SMR) of an AE or ORIO, or expedited review, the IRBMED Regulatory Team will assess the application to determine, to the extent possible, whether the reviewer has a COI. However, it is ultimately the responsibility of the member to self-identify any COI at the time it is known. IRBMED staff will not assign an application to a conflicted IRB Member reviewer. If a previously unreported conflict is identified in the course of reviewing an application, a different reviewer will be assigned to the application.

h) Conflicts of Interest Involving the Institution

Refer to HRPP OM Part 9.IV.

C. IRB Review Policies and Procedures

Refer to HRPP OM Part 3.III.C

1. IRB Jurisdiction and Authority

a) Human Subject Research Studies Reviewed by the IRB

IRBMED reviews studies submitted per the assigned jurisdiction in HRPP OM Part 5.I. Submissions include all materials associated with new project (initial) applications, scheduled continuing review applications, amendments, adverse events (AEs), Other Reportable Information or Occurrences (ORIO) reports, and research that may qualify for exemption. Submissions are routed to the IRBMED office by the PI via eResearch, the web-based system for submission, routing, approval and management of human subject research information.

b) Authority of the IRB to Approve, Disapprove or Require Modification to a Study

Refer to HRPP OM Part 3.III.C

All regulated human subject research conducted by the University must be approved by an IRB or granted exemption by a University IRB;

The IRBMED has the authority to approve, disapprove or require modifications to human subject research under its jurisdiction.

c) Authority of the IRB to Suspend, Terminate or Place Restrictions on a Study

Refer to HRPP OM Part 12.II.D-F

Refer to HRPP OM Part 3.III.C

The IRBMED has the authority to suspend or terminate approval of a study or to place restrictions on the performance of the study. It must document the circumstances under which these actions are taken and make a report to UMOR.

d) Not-Regulated and Exempt Research
The IRBMED requires its staff, consistent with the OHRP Guidance and in consultation with the IRBMED Director or Chairs as appropriate, to make the following determinations with respect to every submission for initial or continuing review:

- That the activity described in the application is “research” as defined in the Common Rule
- If considered research, whether the activity involves “human research” as defined in the Common Rule or “clinical investigation” as defined by the FDA regulations; and
- Whether U-M is engaged in the research and
- Whether the research is exempt from IRBMED oversight.

(1) Not-Regulated

For activities not-regulated as human subject research per HHS and FDA definitions of human subject research, the IRBMED does not require PI to seek a determination of ‘Not Human Subject Research’ from the IRBMED (e.g., review of records preparatory to research, QA/QI, or case studies).

PIs seeking documentation of the not-regulated status may submit an application in eResearch and obtain a system-generated determination letter for qualifying circumstances. The PI may also request that the application may be submitted for IRB review to confirm the circumstances of the not-regulated determination. Determination letters of ‘Not Human Subject Research’ are provided via eResearch.

(2) Exempt

Determination of exemption from 45 CFR 46 may be requested by an investigator via an eResearch application to IRBMED. Exemption may be granted by the IRB Chair or designee, including qualified members of the IRBMED Office staff, or the VPR. Once approved, the research activity is not monitored by the IRB. Assuming the project does not exceed the scope of the assigned exemption, it also is not subject to continuing IRB oversight. Exempt status does not lessen the ethical obligations to human subjects as articulated in the Belmont Report and in disciplinary codes of professional conduct.

By agreement, IRBMED also permits certain qualifying applications for exemption to be reviewed by IRB-HSBS.

e) International Research

Generally, the IRBMED will review all international human subject research projects conducted by U-M investigators under its jurisdiction, rather than deferring review to a collaborating international institution.

(1) Federally Supported

When an international site is engaged in the conduct of a U-M research project and the research is supported by a Common Rule agency, the regulatory requirements of 45 CFR 46 are applied and local IRB or ethics committee review is required. Supporting agencies may require a FWA.

(2) Non-Federally Supported

For international research that is not federally supported, the IRB may apply the same or equivalent protections as those described in the Common Rule and U-M
institutions. IRBMED may require local IRB review, particularly for studies involving more than minimal risk to participants.

Where the international research site is not engaged in the conduct of the research, IRBMED may request a letter of collaboration from an appropriate official agreeing to the conduct of the research.

(3) IRBMED Requirements

Projects conducted in international settings are subject to the same IRB requirements for review and approval of initial applications, scheduled continuing review and review of amendments as projects conducted domestically. A key element of the review process is the assessment of the informed consent process and documents. IRBMED evaluates the informed consent process to ensure that it is culturally sensitive and in a local language that is understandable to the subject, and that the complexity of the information is appropriate for the research population. Informed consent documents and other study materials must be provided to IRBMED in the languages in which they will be offered, as well as in English.

(4) IRBMED Review

IRBMED will consider local research context when reviewing research conducted in international settings. Elements of consideration include:

- Laws and regulations
- Local customs and cultural norms
- Political and socio-economic conditions
- Language and literacy issues

The eResearch application elicits information from the study team regarding their experience with and knowledge of the community and culture in which the research will take place. When IRBMED members do not possess the appropriate cultural knowledge to review research in a particular country or region, IRBMED will seek guidance from consultants with cultural expertise to assist with the review. IRBMED may also request that the investigator seek cultural review by an IRB or ethics committee review or from a government agency in the region. For exempt research, IRBMED does not require documentation of IRB review or other approvals from international sites.

(5) Monitoring

Post approval monitoring, such as project reports to IRBMED by the PI, may be imposed when necessary. As with domestic projects, PIs are obligated to report subject complaints, UaP/UPRISOs and other reports of potential non-compliance to IRBMED. Research subjects are provided with the IRBMED email address and international phone number as part of the consent process.

2. Institutional Approval/Disapproval of IRB Decisions

Refer to IRBMED SOP Part 1.III-IV.

3. Submission of IRB Applications and Reports

Refer to HRPP OM Part 3.III.C.3

The University utilizes eResearch, a web-based system for submission, routing, approval, and management of human research information. eResearch relies upon a role-based
structure that permits only a single PI per application and requires the PI to be the individual that functionally submits the initial, continuing review, amendment, and termination applications within the electronic system. The PI may delegate authority to co-Is or faculty advisors for the submission of AE/ORIO reports. The PI is responsible for the content of each eResearch submission and assumes responsibility for compliance with all regulations, laws, and policies associated with the conduct of the research.

4. General IRB Review and Approval Procedures

a) Determining Whether and Under What Authority the Research is Regulated

Refer to HRPP OM Part 3.III.C.4.a

Beyond the requirements of Common Rule and FDA regulations, the IRBMED staff considers additional regulatory requirements associated with the study design such as HIPAA or required by federal sponsors such as DoD or DoEd. IRB staff utilize guidance documents and reviewer checklist in the eResearch application to ascertain any additional requirements.

b) Reviewing IRB Applications (Initial Applications, Amendments, Scheduled Continuing Reviews (SCRs), and Termination Reports per 45 CFR 46.111 and 21 CFR 56.111 and 21 CFR 50.

Refer to HRPP OM Part 3.III.C.4.b

(1) Information Required for IRB Review

Refer to HRPP OM Part 3.III.C.4.b1

A submission to the IRB that is an initial application, amendment or SCR and regulated by 45 CRF 46 (the Common Rule) or 21 CFR 56 must contain the indicated information.

For initial applications and SCRs, the IRB may request other supporting documentation that, in its discretion, will facilitate a complete and meaningful review of the study, such as sponsor or contract research organization contracts governing the conduct of the research, conflict of interest management plans or FDA documents.

(a) Initial Applications

A PI who intends to initiate a new research study involving human subjects that is subject to IRBMED jurisdiction must submit an initial application in eResearch for IRBMED review and approval. No aspect of the project (including testing performed solely to determine eligibility for the project) may begin until the application has been approved in the eResearch system.

• Description of the professional qualifications of the investigator conducting the research;
• Study Protocols that address:
  o Study title;
  o Purpose of the study;
  o Risks and expected benefits to human subjects;
  o Sponsors of the study and any relevant federal grant applications;
  o Results of previous related research;
  o Participant inclusion/exclusion criteria;
o Justification for use of any special/vulnerable participant populations 
  (e.g., cognitively impaired or children (21CFR50subparts B, C, D);
  o Test article accountability procedures;
• Description of study design (including as needed, a discussion of the 
  appropriateness of research methods);
• Description of interactions and interventions and procedures to be 
  performed, including as applicable, any questionnaires, surveys, or scripts 
  used by PIs or the study team to communicate with participants or their 
  LARs;
• Provisions for managing adverse reactions;
• Copies of the proposed informed consent documents (including all 
  requirements of 21 CFR 50.25(a); including requirements of 21 CFR 
  50.25(b) that are appropriate to the study; meeting all requirements of 21 
  CFR 50.20; translated consent documents, as necessary, considering 
  likely participant population(s); or a request for IRB approval of a waiver of 
  written informed consent;
• A description of the accommodations that will surround the informed 
  consent process, including setting, participant autonomy concerns, 
  language barrier concerns, vulnerable population needs;
• The procedures for documentation of informed consent, including any 
  procedures for obtaining assent from minors; using witnesses, LARs, 
  translators; and the plan for secure storage of informed consent 
  documents;
• Any compensation for injured participants;
• Extra costs to participants for partaking in the study;
• Adequate provisions to protect the privacy of participants (i.e., individually 
  identifiable health information) and to maintain confidentiality (security) of 
  the data;
• Copies of advertisements and any other recruiting materials (including, but 
  not limited to, posters, website contents, videotapes, scripts for telephonic 
  communications), if used;
• An adequate monitoring plan to review data, where appropriate, to ensure 
  the safety of study participants;
• Documentation of approval from other University departments or divisions 
  from which the IRB requests approval or certification that such approval 
  will be obtained before the study begins;
• Documentation of approval, disapproval, or other action from other 
  performance sites (partners) performing the research, if the University 
  (directly or through the PI) has ultimate responsibility for the conduct of the 
  study or performs any coordinating functions including, without limitation, 
  study coordination, recruitment, data management, data storage, 
  monitoring, or otherwise; or certification that such approval will be 
  obtained before the study begins;
• For multi-center trials supported by DHHS, the approved sample informed 
  consent documents and complete DHHS-approved protocol (if any)
• The Investigators Brochure, IND/IDE application or exemption 
  documentation (e.g., an IDE letter), if any, (for studies involving the use of 
  an investigational drug, biologic, or device);
• The clinical case report form (if any);

(b) Scheduled Continuing Reviews (SCRs)
The PI of an approved research study is responsible for submitting an application for scheduled continuing review (SCR) an approval sufficiently in advance of the expiration date of the current approval period to permit IRBMED approval within that period.

The eResearch system generates automatic reminders at 90, 60, and 30 days prior to study expiration. If approval for continuation is not issued prior to the expiration date, the PI must cease all research activity until the IRBMED has issued its approval, with the exception of research-related interventions that are necessary to avoid harm to a subject. Note that expiration of an approval does not constitute a “suspension” of IRBMED approval reportable to OVPR and federal regulations under KHHS, FDA regulation, or these SOPs.

An application for SCR must include at least the following information:

- The number of subjects accrued since the initial application or the previous continuing review application;
- The number of subjects expected to be recruited in the future;
- Any changes in the risk level determination;
- A summary or tabulation of any reports including:
  - unanticipated problems involving risks to subjects or others;
  - subject withdrawals from the project;
  - complaints received along with the resolution;
  - protocol deviations/violations;
  - accidents/incidents involving data, specimens, or facilities;
  - information about risks associated with the research;
  - reports from or to an oversight entity;
  - documentation of any findings made by external inspectors reviewers or auditors such as sponsors, contract research organizations, or government agencies relevant to the conduct of the research project, as well as the PI’s response to the findings or progress reports submitted to study sponsors or the FDA;
  - reportable adverse events, amendments, or modifications made to the research and a summary of pertinent publications/public announcements obtained or discovered (e.g., articles whose findings may have an impact on the IRBMED risk-benefit assessment or on a subject’s willingness to participate in the study);
- A copy of the informed consent documents approved by the IRBMED and currently in use;
- Copies of the FDA-required Annual Reports, which should be uploaded to eResearch for FDA research in which the Principal Investigator holds an IND or IDE necessary for the study.
- These materials provide the primary reviewer and IRBMED members with the relevant information necessary to determine whether the study continues to meet the regulatory criteria for approval.

(c) Information Required for Study Closure (Termination)

The PI of an approved project is expected to notify the IRBMED upon completion (and/or termination) of a study. A study termination report should include at least the following information:

- Affirmation that the involvement of human subjects and use of identifiable
human data or specimens in research has concluded

- Description of the plan for secure storage of data and indication whether data will be de-identified
- Number of subjects enrolled in the study
- Number of subjects completing the study
- Number of subjects that withdrew from the study and the reasons for withdrawal
- Number of subject complaints about the project and description/resolution of those complaints
- Number of AEs reported to IRBMED during the study (including any reported concurrently with submission of the termination report)
- For research governed by the FDA, the final report that is submitted to the FDA by the IND/IDE sponsor

(2) Review Process / Primary Reviewers

Refer to HRPP OM Part 3.III.C.4.b.2

(a) Review Process

IRBMED Regulatory Teams and primary reviewers must receive sufficient information (IRBMED SOP Part 3.III.C.4.b) prior to review of submissions to prepare their recommendations for approval of the research.

IRBMED Regulatory teams and primary reviewers will assess and review the eResearch submission and all other supporting documentation depending on the submission type, to satisfy requirements for completeness, consistency, and compliance with University policy, 45 CFR 46.111 (the Common Rule), FDA regulatory requirements (21 CFR-56.111), or other regulatory rules or regulations (HIPAA Privacy Rule, HITECH Security Rule FERPA or other federal, oversight activities) prior to presentation for board review. (Refer to IRBMED Part 3.II.C.4.b).

An IRBMED Regulatory Team will be assigned to each IRB submission in eResearch. In general, research submitted by the PI for review via eResearch is assigned to the IRBMED Regulatory Team overseeing the department (e.g., therapeutic area) or type of research being conducted; for example, a clinical trial submitted by an PI in the Department of Orthopedics is generally assigned to the IRBMED Regulatory Team overseeing that department’s research as part of their review portfolio. Additional guidance is available to address sharing submissions among Regulatory Teams for purposes of backup.

To facilitate the review process, the IRBMED Regulatory Team (or designee) may request clarification or revisions to any or all of the application documents prior to sending to a primary reviewer. Upon completion of their review, the Regulatory Team forwards their Staff Reviewer checklist and comments for primary review to:

- A single member qualified and designated for expedited review (Refer to HRPP OM Part 3.III.C.5); or
- A primary reviewer, with relevant scientific expertise, who will present their review to a convened board

Following IRBMED Regulatory Team review and at the time most identified
issues have been resolved with the study team, the IRBMED Regulatory Team will select an IRB member to function as the “primary reviewer” based on, but not limited to, the member’s expertise, experience, and/or representation of pertinent subject population, as indicated on the member’s Curriculum Vitae, documentation of community experience, or responses to the periodic IRBMED survey of expertise.

If the submission qualifies for expedited review the primary reviewer must also have the appropriate education and experience as determined by the IRBMED to be designated as an expedited reviewer. Refer to IRBMED SOP Part 3.III.C.5 for the expedited review procedure.

The primary reviewer:

- Assesses the initial application, amendment or SCR, together with, ICDs, and all supplemental materials (including, if applicable, the grant application, protocol, recruitment materials, etc. and the elements detailed in IRBMED SOP Part 3.III.4.b).
- Documents his/her review in the eResearch Reviewer Checklist of the application before the convened board meeting where it will be presented;
- May contact the PI in advance of the board meeting for additional information or clarification;
- Leads the discussion of the application under review at the convened meeting;
- May not have a COI regarding the project under review and is expected to notify the IRB staff and Chair of any conflict at the time the review assignment is offered or if previously unanticipated conflict arises during the review.

Compensation to subjects, if any will be in accordance with University policy. The IRBs will review payment arrangements offered to subjects. Their review will ensure the following:

- The amount of payment, the proposed information collected, and the method and timing of disbursement neither is coercive nor presents undue influence or places the subject at risk.
- Where appropriate, credit for payment accrues as the study progresses and may not be contingent upon the subject completing the entire study.
- Any amount paid as a bonus for completion is reasonable and not so large as to unduly induce subjects to stay in the study when they would otherwise have withdrawn.

All information concerning payment, including the amount and schedule of payments, is set forth in the consent document.

No aspect of a study (including review of medical records performed solely to determine eligibility for the study) may begin until the submission has been approved in eResearch by the IRBMED.

In some instances, the IRBMED Regulatory Team, after consultation with the PI and/or study team, may enter changes into the eResearch application for the purposes of assisting the study team and facilitation of the review process. This process, designated as Staff Edit Rights (SER) is described in detail,
including the types of changes that are authorized in an additional guidance document.

The IRBMED Regulatory Team may consult with advisory units (i.e., the MICHIND/IDE Assistance Program [MIAP]) for preliminary review and assistance with research that may require an Investigational New Drug (IND) or Investigational Device Exemption (IDE) application. (Refer to IRBMED SOP Part 2.II)

(b) Timing of Distribution of Materials

The IRBMED Regulatory Teams assign the applications to the eResearch meeting agenda on a rolling basis until the agenda is full. Adjustments may be made to the final agenda to accommodate any reviews that are time-sensitive in nature. Distribution of application materials to the Primary Reviewer and board members generally occur not later than 4 business days before the meeting unless a time-sensitive submission is added after that date.

A secondary reviewer may be assigned if additional expertise is deemed necessary. The secondary reviewer may be another voting member of the Board or a non-voting member/consultant to the Board. All study documents will be made available to the secondary reviewer.

(c) Regulatory Criteria for Board Review

Initial Applications and Amendments

All eResearch applications are first reviewed by qualified IRBMED Regulatory staff to assure that the application is complete, all relevant materials are provided, and that the investigator has addressed all necessary regulatory criteria. The application is then assigned for review by experienced IRBMED Board Member(s) to determine that the criteria in 45 CFR 46.111 or 21 CFR 56.11 are met and that the study design is adequate to protect the subjects from increased risk and yield expected knowledge. This includes examination of the following:

- Risks to subjects are minimized
  - by using procedures that are consistent with sound research design and do not unnecessarily expose subjects to risk;
  - whenever appropriate, by using procedures already being performed on the subjects for diagnostic or treatment purposes; and
  - when adequate resources are available to protect and minimize harm to human subjects.

- Sound Research Design / Scientific Review
  - Risks to subjects are reasonable in relation to anticipated benefits, if any, to subjects, and the importance of the knowledge that may reasonably be expected to result;
  - Selection of subjects is equitable;
  - When appropriate, the research plan has an adequate data and safety monitoring plan; (Refer to IRBMED SOP Part 3.III.C.6.f)
  - There are adequate provisions to protect the privacy of subjects and to maintain the confidentiality of data (Refer to IRBMED SOP Part 3.III.C.6.g);
  - When appropriate, additional safeguard have been included in the study to protect the rights and welfare of vulnerable subjects.
(Refer to IRBMED SOP Part 3.III.C.6.h);

- Studies that have received peer or scientific review indicate in the eResearch application the name of the unit or person(s) who performed the review;
- For student applications it is expected that the faculty advisor has reviewed the study for scientific merit before it is submitted to IRBMED;
- For studies that receive federal support (and thus a scientific review) the grant application must be uploaded into the eResearch system;
- For studies conducted in the Comprehensive Cancer Center, all studies are reviewed by the Protocol Review Committee (as a Core Committee of eResearch).

- Risks to subjects are reasonable in relation to anticipated benefits, if any, to subjects, and the importance of the knowledge that may reasonably be expected to result.
- Assessment of risks and benefits of the research will include consideration of immediate medical as well as societal benefit. Refer to IRBMED Guidance – Risk Guidelines for using Magnitude of Harm in Categorizing Risk Level.
- In evaluating risks and benefits, the IRBMED will consider risks and benefits that may result from the research. The IRBMED does not consider possible long-range effects of applying knowledge gained in the research (i.e., the possible effects of the research on public policy) to be among those research risks that fall within the purview of its responsibility.
- Selection of subjects for participation in the project is equitable. In making this assessment, the IRBMED takes into account the characteristics of the subject population, the purposes of the research, the setting in which it will be conducted, recruiting methods and materials, and other relevant information.
- Informed consent (unless waived) will be sought from prospective subjects or their legally authorized representatives before enrollment in the protocol, in a manner that minimizes the likelihood of coercion or undue influence and will be documented on a form approved by the IRBMED (per 46.111[a][5]). Refer to IRBMED Informed Consent and Assent Templates.
- When appropriate, the research plan makes adequate provision for monitoring the data collected to ensure the safety of subjects.
- When appropriate, there are adequate provisions to protect the privacy of subjects and to maintain the confidentiality of data.
- When some or all of the subjects are likely to be vulnerable to coercion or undue influence (such as children, prisoners, pregnant women and fetuses, handicapped, mentally disabled persons or economically or educationally disadvantaged persons), additional safeguards have been included in the study to protect the rights and welfare of these subjects (e.g., to minimize risks peculiar to these groups and the possibility of coercion or undue influence).

Scheduled Continuing Reviews (including Terminations)
The IRBMED conducts scheduled continuing review of any research study
subject to its oversight at intervals appropriate to the magnitude of risk of the study and other considerations, but not less than once each year (except for research meeting the criteria for regulatory flexibility). The IRBMED will:

- Assess the current risk level of the project and, if necessary, revise the risk level (decrease or elevate) commensurate with the activity being conducted; and
- Require any other changes warranted in accordance with the changes in risk level.

Termination of a study, whether due to completion or other reason, is submitted via the SCR mechanism. For further information about termination reports refer to IRBMED SOP Part 3.III.4.b.1.c.

(d) Board Actions

The IRB may vote to take any of the following actions with respect to an application for initial, amended, or scheduled continuing review:

(i) Approve the Submission as Presented to the Review Board

Submissions will be eligible for approval only if the criteria listed IRBMED SOP Part 3.III.C.6 are met.

(ii) Approve the Submission with Contingencies

Approval will be contingent on specified changes to the protocol, ICDs or other items that will be made by the PI prior to initiating the research.

- Approved Pending Office (APO)

Approval of the submission is contingent on specified administrative changes requested and clearly defined by the IRB. The requested changes, such as correction of typographical errors, specified wording changes, or other items will be made by the PI or study team.

- Approved Pending Reviewer (APR)

Approval of the submission is contingent on specified changes requested by the board. If there are contingencies remaining and they are of an administrative nature as clearly defined by the Board (e.g., correction of typographical errors or specified wording changes to an informed consent) the reviewer may request that the IRBMED Regulatory Team verify those changes. Once all contingencies are met and they meet with the reviewer’s approval (and that of the designated IRBMED Regulatory Team approval may be granted.

If the PI disagrees with the IRBMED request or proposes an alternate change, the approval status of the application will be “deferred,” and the application must be re-presented at a subsequent board meeting in order to obtain approval, unless the application under the regulations qualifies for review in an expedited fashion.

The date of the vote for approval as APO or APR shall be deemed the date of approval, regardless of when the specified changes are made by the PI and submitted to the IRBMED. The IRBMED may, in its discretion, require that the PI respond to required changes within a specified period and instruct that if the response is not received, the
application will be considered withdrawn or reassigned to a board action deferred status.

(iii) Board Action Deferred (BAD)

In the event that a submission requires changes that are significant or substantively require more than simple concurrence of the PI, action on the submission shall be deferred.

Board action may be deferred on any submission without a vote. If neither a motion to approve nor a motion to disapprove is carried, the action is automatically deferred. In this case, the PI may be instructed to submit additional information or revisions required by the IRBMED before reconsideration of the submission. The IRBMED may, in its discretion, require that the PI respond within a specified period and instruct that if the response is not received, the application will be considered withdrawn.

(iv) Disapproval of the Application

Refer to IRBMED SOP Part 3.III.C.4.b.1.c

PIs will be notified of the reasons for disapproval and afforded the opportunity to appeal the decision.

(v) Suspension or Termination of IRBMED Approval

Refer to HRPP OM Part 12.III.

The IRBMED also may suspend or terminate approval of research that it determines, after appropriate review and deliberation:

- is not being conducted in accordance with IRBMED requirements;
- has been associated with unexpected harm to subjects; and/or
- cannot minimize risks to subjects or maintain a favorable risk-benefit balance. Any suspension or termination of approval under this provision shall include a statement of the reasons for the action and inform the PI of institutional notification and reporting requirements.

Suspension of Research Activity

Suspension is the temporary closing of a human subject research project or discontinuing a PI's privilege to conduct human subject research. The suspension may be partial, in that certain activities may continue while others may stop; or it may be complete, in that no activity related to the research may proceed.

Termination of Research Activity

Termination is the ending of all activities related to human subject research or a PI’s privilege of conducting human subject research except for the continuation of follow-up activities necessary to protect human subject safety.

Refer to IRBMED SOP Part 12 for reporting a suspension or termination of IRBMED approval.

(3) Timeliness of Submissions and Reviews
(a) Notices of Expiration

It is the PI’s responsibility to submit an application for continuing review (SCR) before expiration of IRBMED approval and in ample time for IRBMED review.

- eResearch-generated reminder notices are sent to PIs and designated study team members at 90, 60 and 30 days prior to the expiration date of the current approval period.
- If IRBMED has not reviewed and approved the SCR application by the expiration date of the current approval (regardless of the reason or circumstances), the study will be considered lapsed and the research must stop unless the IRBMED reviews, approves, and document that it is in the best interest of individual subjects currently participating in the study to continue the research interventions or interactions;
- Enrollment of new subjects or intervention and interaction with subjects during a lapse is prohibited, as is data analysis.
- Sponsored project resources (e.g., government or private) must not be expended for unallowable activities;

(b) Lapses in Approval and Administrative Termination

If an approved research project is not renewed or terminated within three months after the date of previous approval expiration, the IRBMED may consider the research to have been completed or discontinued, and may administratively terminate that protocol notwithstanding the lack of a study completion or termination report.

- Notification will be sent to the PI prior to termination; the IRB will consider evidence from the PI in the event the PI wishes to submit a continuing review. The IRB may determine that submission of a new application is necessary, rather than renewing approval of the now-lapsed application.
- An administrative termination under this provision does not constitute a suspension or termination of IRBMED approval reportable to UMOR and regulatory agencies under 45 CFR 46.113 or these SOPs.

For projects reviewed, but not approved, by the IRBMED due to outstanding contingencies, the IRBMED staff may administratively withdraw the project after notification to the PI. Withdrawals of applications by IRBMED staff may occur after abandonment of an application or communication with the PI of their intent to modify their plans not to finalize the contingencies.

Additional guidance is available regarding Administrative Terminations and Withdrawals.

(4) Notice and Appeal of IRB Determinations

(a) Notification of Determinations

Following an IRB meeting, the IRBMED Regulatory Team shall prepare electronic notification to inform the PI of each submission upon which a vote was taken, and on the outcome of the vote. The notification shall include at least the following information:
- The IRB’s decision and date it was reached;
- For an approved submission, the approval expiration date and notification...
of any interim reporting requirements;
• A list of currently approved documents, e.g., the informed consent and protocol with specific reference to version number as applicable;
• For a project approved contingent on specified changes to be made to the protocol, ICDs, or otherwise, a description of the specific modifications necessary to secure approval.
• The IRBMED may, in its discretion, require that the PI respond to required changes within a specified period and instruct that if the response is not received, the application will be considered withdrawn or reassigned to deferred status.
• For a disapproved, suspended, or terminated project, the reasons for the IRB’s decision and notification of the PI’s right to respond in person or in writing.

Documentation of all IRBMED determinations shall be available for review by the Medical School Associate Dean for Regulatory Affairs, UMOR, IRBMED members, and authorized consultants.

A copy of any notification of a board suspension or termination of a project shall be delivered under cover letter to UMOR for further disposition and notification to other interested parties, as necessary, such as government authorities with jurisdiction (i.e., the FDA and OHRP) and, in the case of a sponsored project, ORSP.

The IRBMED may, in its discretion, report disapprovals, or other actions to UMOR as it deems necessary or appropriate.

(b) Appeal of Determinations

The PI may appeal any decision by the board through a telephonic or written (e.g., email) request to the Co-Chairs or Vice-Chairs of the reviewing board.

c) Frequency of Review

In general, the approval period for an initial research application begins on the date it is approved by the IRBMED and expires 364 days later, which is the last date of the approval period. For example, an application will have an approval date of 9/30/15 and an expiration date of 9/29/16.

The IRBMED may approve an initial application or SCR for intervals of less than one year when warranted. Criteria for this consideration include, but are not limited to:

• The overall risk of the study, with the highest risk studies reviewed more frequently;
• Data safety monitoring plan requirements;
• Demonstrated the need for additional oversight of the PI and study team;
• Questions about sufficiency of the data to lead to generalizable knowledge;
• Excessive numbers of serious adverse events (SAEs) or protocol deviations;
• The protocol is subject to complex regulatory compliance requirements, such as research involving investigator-held IND or IDE;
• The research is being conducted in an off-site location(s) and the IRBMED is serving as the IRB-of-record;
• An investigator conducting the research has a potential COI that warrants more frequent reporting and review.

There may be additional circumstances that the IRB would consider as significant to warrant the additional oversight.

The University permits IRBs to undertake flexibility or demonstration projects that may lengthen an approval period beyond one year.

*Refer to IRBMED SOP Part 3.III.C.4.c*

d) Monitoring and Verification by IRB

*Refer to IRBMED SOP Part 12.II.G*

The IRBMED is responsible for overseeing the safety of human research subjects and has the authority to suspend or terminate human subject research that:

• is not being conducted in accordance with federal and IRBMED requirements (45 CFR 46.103(b)(5)), 45 CFR 46.113, 21 CFR 56.113 and IRBMED SOP Part 12); and/or
• has been associated with unexpected serious harm to human participants in research.

The IRBMED may, at its discretion,

• Perform monitoring of studies both for-cause (e.g., alleged noncompliance) and not-for-cause (e.g., random review for quality assurance purposes);
• Request monitoring from the Office of Human Research Compliance Review (OHRCR) – via UMOR– of a study; in addition to information received through the initial application, any amendments, annual SCRs, and analyses of interim reports, such as AEs and audit reports. For example, the IRBMED may choose to undertake extra monitoring for research which presents greater than minimal risk, or to gauge the progress of recruitment of vulnerable participants, or to follow the research progress on controversial subject matter;
• Consider the frequency and nature of AEs reported to date.

Criteria for monitoring may include, but is not limited to, the following:

• Complex projects involving unusual levels or types of risk to subjects;
• Projects conducted by PIs who previously have failed to comply with applicable regulations, institutional or IRBMED requirements;
• Projects where other concerns about possible material changes occurring without IRBMED approval have been raised (e.g., major changes to the study protocol were made without an amendment);
• Projects involving vulnerable populations;
• Complaints received regarding the study.

The IRBMED may also choose to monitor one or more of the projects of a single PI in consideration of the experience of the PI or as follow-up to previous reports of complaints, non-compliance, or prior IRBMED interactions with the individual. Monitoring may include, but is not limited to, providing the IRBMED copies of or access to any or all of the following:

• Signed informed consent documents;
• Study files and research records;
• Drug dispensing/IDS logs;
• Subject records;
• Lab test procedures, results and raw data;
• Observation of study activity (e.g., witnessing the informed consent process);
• Review of study by an outside auditor;
• Interviews of study personnel;
• Interviews of research subjects;
• Site visits to research locations;
• Monitoring reports/findings;
• Independent third party monitoring reports;
• Projects involving vulnerable populations;
• Reports by the Data and Safety Monitoring Board (DSMB).

e) Reporting Changes in Research to IRBMED (Amendments)

All amendments to research must continue to meet the requirements of 45 CFR 46.111 or 21 CFR 56.111 in order to be approved.

Once a project has been approved a PI may not make any changes to the project (e.g., changes to the protocol, ICD, recruitment materials or subject incentive) without prior IRBMED review and approval, unless necessary to eliminate apparent immediate hazards to the subjects. Any change made without prior approval to avoid a hazard must be reported promptly to the IRBMED.

The IRBMED will scrutinize any proposed amendments to determine the degree to which risks to human subjects may have changed, whether there is any need to revise the ICDs or informed consent process, whether proposed changes in the ICD are appropriate, and/or whether there is any need to notify previously enrolled subjects of the changes and if reconsenting of the subjects is necessary.

Reportable changes may include, but are not limited to:
• Proposed changes in risks or benefits to subjects;
• Proposed amendments to the study protocol, including changes to the eligibility criteria, recruitment materials, questionnaires, surveys, scripts and subject payments;
• Proposed amendments to the Investigators Brochure or equivalent documentation;
• Proposed amendments to previously approved ICDs;
• Proposed changes in Investigators (including PIs, Co-Is, researchers) or performance sites;
• Proposed changes to subject population;
• Proposed changes in any other aspect of the research.

At the request of a PI, the IRBMED Chairs or IRB will consider or agree to acknowledge a voluntary hold on subject enrollment or delay any portion of research activities to facilitate significant changes to a research study and further IRBMED review of the study or its conduct.

The date of IRBMED approval of an amendment does not extend the approval period of the study.

f) Preventing Lapses in IRB Approval

Refer to IRBMED SOP Part 3.III.C.4

5. Expedited Review
Refer to HRPP OM Part 3.III.C.5.

DHHS regulations at 45 CFR 46.108(b) and FDA regulations at 21 CFR 56.08(c)(i) identify certain types of research that may be reviewed and approved by “expedited review.” The following types of research may be considered for an expedited review process:

- The research involves no more than minimal risk to subjects;
- The identification of the participants on their responses will not reasonably place them at risk of criminal or civil liability or be damaging to their financial standing, employability, insurability, reputation, or be stigmatizing, unless reasonable and appropriate protections will be implemented so that risks related to invasion of privacy and breach of confidentiality are not greater than minimal;
- The research is not classified; and
- The research falls into one of more of the categories of projects or applications appearing on a list of expeditable studies published by the Secretary of the Department of Health and Human Services, and only in those categories (see 45CFR46.110 and 21CFR56.110).
- Minor changes in previously approved research during the period (of one year or less) for which approval is authorized.

Under an expedited review procedure, the IRBMED Chairs, or an experienced IRBMED reviewer designated by the Chair from among the members of the IRBMED reviews the research submission (refer to IRBMED SOP Part 3.III.B.7.g). Consultants may assist the IRBMED in the review of issues which require expertise beyond, or in addition to, that available with current full or alternate members of the IRBMED boards.

The research submitted for expedited review requires the same materials that a convened board would receive for standard submission.

The IRBMED Regulatory Team performs a regulatory review of the application followed by assignment to an expediting reviewer. Following reviewer assignment, and at the reviewer’s discretion, submissions eligible for expedited review may be referred to the full IRBMED for a discussion and vote.

When applicable, questions or requirements pertaining to an expedited submission will be communicated to the PI by the IRBMED Regulatory Team or the expediting reviewer and must be addressed to their satisfaction prior to approval of the submission. The IRBMED Regulatory Team and expediting reviewer will document findings, determination, or recommendations on the Reviewer Checklists in eResearch.

a) Expedited Review Categories of Research

The interactions and interventions in the research must present no more than minimal risk to subjects. The regulatory definition of minimal risk is that “the probability and magnitude of harm or discomfort to participants that is anticipated in the research are not greater in and of themselves than those ordinarily encountered in daily life or during the performance of routine physical or psychological examination or tests”.

(1) The research conducted must meet the list of research categories as defined in UM’s HRPP OM –Categories of Research and cited below: The categories apply regardless of the age of subjects, except where noted.

Clinical studies of drugs and medical devices only when one of the following conditions is met:
(a) Research on drugs for which an IND (21CFR312) is not required or 
(b) Research on medical devices for which an IDE application (21CFR812) is not 
required; or the medical device is cleared/approved for marketing and the 
medical device is being used in accordance with its cleared/approved 
labeling.

(2) Collection of blood samples by finger stick, heel stick, ear stick or venipuncture as 
follows:

(a) From healthy, non-pregnant adults who weigh at least 110 pounds. For these 
subjects, the amounts drawn may not exceed 550 ml in an 8 week period 
and collection may not occur more frequently than 2 times per week; or  
(b) From other adults and children (for definition see 45CFR46.402(a)), 
considering age, weight and health, the collection procedure, the amount of 
blood to be collected, and the frequency with which it will be collected. For 
these subjects, the amount drawn may not exceed the lesser of 50ml or 
3ml/kg in an 8 week period and collection may not occur more frequently 
than 2 times per week.

(3) Prospective collection of biological specimens for research purposes by 
noninvasive means (e.g. hair and nail clippings, teeth in need of extraction, saliva, 
excreta, dental plaque and calculus, mucosal and skin cells, sputum);

(4) Collection of data through noninvasive procedures (not involving general 
anesthesia or sedation) routinely employed in clinical practice, excluding 
procedures involving x-rays or microwaves; where medical devices are used they 
must be cleared/approved for marketing (but not for new indications).

(5) Research involving materials (data, documents, records, or specimens) that have 
been collected, or will be collected solely for non-research purposes (such as 
medical treatment or diagnosis). Refer to 45 CFR 46.101(b)(4) for some 
examples in this category that are exempt).

(6) Collection of data from voice, video, digital, or image recordings made for 
research purposes.

(7) Research on individual or group characteristics or behavior (including, but not 
limited to research on perception, cognition, motivation, identity, language, 
cultural beliefs, social behavior) or research employing survey, interview, oral 
history, focus group, human factors evaluation or quality assurance 
methodologies. Refer to 45 CFR 46.101(b)(2) and (b)(3) for some examples in 
this category that are exempt.

(8) Continuing review of research previously approved by a convened IRB as follows:

(a) Where the research is permanently closed to the enrollment of new subjects; 
all subjects have completed all research-related interventions; and the 
research remains active only for long-term follow up of subjects; or  
(b) Where no subjects have been enrolled and no additional risks have been 
identified; or
(c) Where the remaining research activities are limited to data analysis

(9) Continuing review of research, not conducted under an investigational new drug application or investigational device exemption where categories two (ii) through eight (viii) do not apply but the IRB has determined and documented at a convened meeting that the research involves no greater than minimal risk and not additional risks have been identified.

Additions to, and extrapolation from, this list by the institution or the IRBMED are not appropriate. For example, it is inappropriate to use an expedited review procedure for the initial review of research that involves minimal risk but does not appear in the categories of research published in the Federal Register or for research that involves greater than minimal risk.

b) Expedited Review of Minor Changes

The IRB also may use expedited procedures to review minor changes in previously approved research during a period for which approval is authorized. For purposes of this policy, a proposed change in research is deemed "minor" if it does not significantly affect an assessment of the risks and benefits of the study and does not substantially change the aims or design of the study. A modification cannot be deemed minor if it involves the addition of procedures that involve more than minimal risk or does not fall into federal categories (1) – (7) of research that can be reviewed by expedited procedures.

Examples of minor changes to a research study include, but are not limited to:

• Addition or deletion of study team members;
• Addition of procedures that do not significantly increase risk to subjects, considering the original purpose and study design of the approved study (i.e., new procedures that fall under any of the expedited categories can usually qualify as minimal risk);
• Removal of research procedures that would thereby reduce the risk to no more than minimal (i.e., procedures now meet expedited research categories);
• Addition of non-sensitive questions to unvalidated survey or interview procedures;
• Addition of or revision to recruitment materials or strategies;
• Changes to improve the clarity of statements or to correct typographical errors, provided that such changes do not alter the content or intent of the statement.

c) Expedited Reviewers

Generally, IRB Chairs appoint experienced IRB members to serve as expediting reviewers. For purposes of this policy, a member is deemed experienced if he or she has completed all mandatory education for IRB members, has served on the IRB for a minimum of six (6) months or has described and documented appropriate experience, and has been approved by the IRB chairs as qualified to perform expedited reviews.

d) Expedited Review Determinations

In conducting expedited review, the IRBMED reviewers may exercise all of the authorities of the IRBMED, except that they may not disapprove the research, in accordance with the non-expedited review procedure set forth in 45 CFR 46.108(b)
and 21 CFR 56.108(c)(i). The reviewer may either approve, require modifications (to secure approval), or refer the research to the convened IRBMED for review (for example, if they determine the study has a change in risk level due to a change in the protocol).

The IRBMED Regulatory Team will prepare notification of any expedited determinations that will be provided electronically to the appropriate IRB. On a monthly, but not less than quarterly basis, a list of all expedited review approvals inclusive to that period will be provided on an IRB meeting agenda for acknowledgement at a convened IRB meeting.

The notification shall include at least the following information:

- the reviewer’s name;
- the submission title and study number;
- a description of the qualifying research category; and
- the expediting reviewer’s decision and the date it was reached.

For an approved project, the notification will also include:

- the approved expiration date; and
- notification of any interim reporting requirements.

For a project approved contingent on specified changes being made to the protocol, ICDs, or otherwise, the notification will include a description of the specific modifications necessary to secure approval.

The IRBMED may, at its discretion, require that the PI respond to required changes within a specified period and instruct that, if the response is not received, the application will be considered withdrawn or reassigned to deferred status. The PI may appeal any decision by the board per procedures outlined in IRBMED SOP Part 3.III.C.4.b.4.b.

e) Limitations of Use of Expedited Review

The expedited review procedure may **not** be used where:

- identification of the subjects and/or their responses would reasonably place them at risk of criminal or civil liability;
- would be damaging to the subjects’ financial standing, employability, insurability, or reputation; and/or
- would be stigmatizing, unless reasonable and appropriate protections are implemented so that risks related to invasion of privacy and breach of confidentiality are no greater than minimal.;
- research is classified; and
- research involves prisoners.

Other limitations may be placed on expeditable research by an IRB Chair, the VPR, or, for research that is federally supported or FDA-regulated, the relevant department or agency head per HRPP OM Part 3.III.C.5.

f) Demonstration/Flexibility Projects

Research demonstration/flexibility projects allow for the addition of expedited review
categories for research that are not federally sponsored or FDA regulated and meet other qualifying criteria. Refer to Innovation and Demonstration website.

6. Criteria for IRBMED Approval

Regulations at 45 CFR 46.111 (Common Rule) or 21 CFR 56.111 (FDA-regulated research) delineate specific criteria for the approval of research. The IRBMED shall determine that all of the following requirements are satisfied before approving proposed research:

a) Scientific Merit and Feasibility

Refer to HRPP OM Part 3.III.C.6.a.

The IRBMED reviews all initial protocols for scientific merit and feasibility and considers supporting background scientific information. Scientific merit is examined in relationship with the risks and benefits of the research to human participants. The Protocol Review Committee (PRC) reviews all UM Comprehensive Cancer Center protocols prior to IRBMED review and approval.

When performing the scientific review using the eResearch Reviewer Checklist, the primary reviewer shall ascertain and indicate that each of the listed elements is adequately addressed. The primary reviewer may also add additional comments and provide specific information regarding any scientific shortcomings identified in the proposal.

No protocol may be approved unless its scientific validity has been ascertained and documented using the Reviewer Checklist.

b) Minimizing Risk: 45 CFR 46.111(a)(1)

Refer to HRPP OM Part 3.III.C.6.b.

To approve research, the IRBMED must determine that risks are minimized by using procedures that are consistent with sound research design and do not expose participants to unnecessary risks. Where appropriate, the research project design should include procedures that are already used with the subjects for diagnostic or treatment purposes.

The IRBMED verifies that the research plan, including research design and methodology, will not place participants at unnecessary risk. This includes the risk that the research is inappropriately designed or is lacking in statistical power, such that meaningful results cannot be obtained. To assist the IRBMED staff with making these determinations, the eResearch application provides guidance materials, including checklists.

The IRBMED shall also consider the professional qualifications of the research team, as well as the resources available to the research team at the specific location(s) where the research will be conducted, including but not limited to facility resources such as the testing and safety equipment. PIs and Co-Is are expected to maintain appropriate professional credentials and licensing privileges.

c) Risk-Benefit Analysis

Refer to HRPP OM Part 3.III.C.6.c.

All research studies, regardless of the type of review (initial or SCR; convened board or expedited), undergo a risk/benefit assessment.

The IRBMED will review the eResearch application to evaluate the risks versus
benefits of the study, using supporting documents, scientific references, IRBMED Regulatory Team and primary reviewer Checklists, and recommendations provided by consultants (as appropriate).

The initial step in evaluating a study for risk is to determine if the study meets the federal regulatory definition of minimal risk, i.e., “the probability and magnitude of harms or discomfort anticipated in the research are not greater in and of themselves than those ordinarily encountered in daily life or during the performance of routine physical or psychological examinations or tests (45 CFR 46.102[i]).” Note: Prisoner research utilizes a different definition of minimal risk (45 CFR 46.303[d]).

In determining whether a study presents no more than minimal risk to the subjects, the IRBMED considers the following:

- The PI’s assessment of the subjects’ risk level as presented in the eResearch submission;
- Whether the study procedures are consistent with sound research design;
- An evaluation of the probability (likelihood) of harm occurring and the magnitude (potential severity) of possible harm;
- An evaluation of whether the subjects are vulnerable in some way;
- An evaluation of the steps taken, or planned, by the PI to alleviate the potential harms (including the quality of the data safety monitoring plan, as appropriate);
- The PI’s history of compliance with research protocols and IRBMED procedures.

Generally, studies with a low probability of harm are considered no more than minimal risk. If the study does not meet the federal definition of minimal risk, then IRBMED evaluates the design of a proposed study to ensure that:

- It is consistent with fulfilling its scientific mission;
- risks are minimized; and
- potential benefits of the research are maximized as much as possible within the confines of the research study.

Refer to IRBMED guidance: Guidelines for Using Magnitude of Harm in Categorizing Risk Level on the IRB website.

The IRBMED does not consider possible long-range effects of applying knowledge gained in the research (e.g., the possible effects of the research on public policy) to be among those research risks that fall within the purview of its responsibility.

The IRBMED will rely on the expertise of its membership to evaluate the risks and benefits of a research proposal. Alternatively, if physical risks are difficult to assess or are outside the scope of expertise of IRBMED, the protocol may be referred to another IRB according to the policies outlined in the HRPP OM Part 5.I.

d) Equitable Subject Selection: 45 CFR 46.111(a)(3)

Refer to HRPP OM Part 3.III.C.6.d

To approve research, the IRBMED must determine that the selection of participants is equitable. This reflects UM’s adherence to the Belmont Report’s concept of “Justice”. In making this determination, the IRBMED will evaluate:

- the characteristics of the subject population;
- the purposes of the research;
- the setting in which it will be conducted;
- the recruiting methods and materials used; and
- the subject inclusion/exclusion criteria.

The IRBMED should be especially cognizant of special considerations for research involving vulnerable participant populations such as, but not limited to neonates, children, prisoners, pregnant women and fetuses, cognitively impaired persons, or economically or disadvantaged persons. Generally, a population that stands no chance of benefitting from the research should not be selected to assume the risk.

The IRBMED should be mindful of the importance of including members of minority groups in research, particularly when the research holds out the prospect of benefit to individual subjects or the groups to which they belong. Non-English-speaking subjects should not be systematically excluded because of inconvenience in translating ICDs. The IRBMED should also ensure that subjects are not selected from only one group of people simply because it is convenient.

The IRBMED should be mindful of the desirability of including both women and men as research subjects and should not arbitrarily exclude the participation of persons of reproductive age. Exclusion of such persons must be fully justified and based on sound scientific rationale.

e) Informed Consent and Parental Permission

Refer to IRBMED SOP Part 11.III.A

Refer to IRBMED Guidance on Informed Consent on IRBMED website.

(1) General Requirements

Refer to IRBMED SOP Part 11

Refer to IRBMED Guidance on Informed Consent

Comprehensive informed consent requirements and application of those requirements are provided in IRBMED SOP Part 11 and on the IRBMED website. Additional guidance on the website includes links to regulations, templates and suggested wording on the IRBMED website. Throughout this section the term “consent” refers to both “consent” and “parental permission”.

Informed consent (IC) will be sought from each prospective subject or the subject’s legally authorized representative (LAR), in accordance with, and to the extent required by 45 CFR 46.116 and 21 CFR 50.20 Subpart B. Except as otherwise approved by the IRB or allowed under FDA regulations (21 CFR 50.23 Subpart B), no PI may involve a human subject in research unless the PI has obtained the legally effective informed consent of the subject or the subject’s legally authorized representative (LAR).

The PI will submit ICDs for IRBMED review (including written ICDs, oral scripts), descriptions of the process to obtain informed consent from subjects, and any requests for waiver(s) or alteration of informed consent or waiver(s) of documentation of informed consent, in the eResearch submission to the IRB. IRBMED will review the proposed informed consent process, including ICDs for each submitted application to assure that subjects or their LARs provide legally effective, voluntary, informed consent.
In its review of ICDs, IRBMED will ensure that all required elements of consent as well as any additional elements, as appropriate, are included per 45 CFR 46.116. It will also ensure that the documents do not contain any exculpatory statements suggesting that any of the subjects’ legal rights are being waived, or that the PI, sponsor, or U-M may be released from liability for negligence.

The IRBMED will assess applications and issue waivers of documentation or waivers of some or all of the elements of informed consent, where appropriate under regulatory guidance.

Except as otherwise approved by the IRBMED or allowed under FDA regulations, no PI may involve a human subject in research unless the PI has obtained the legally effective informed consent of the subject or the subject’s LAR.

The IRBMED will evaluate the plans for obtaining consent by confirming the following:

- The consent process is facilitated by a person knowledgeable about the study, its enrollment criteria, its risks and benefits, and alternatives to participating in research (usually a PI or Co-I, although other study team members, for example, a research study coordinator or research nurse, may also be qualified and designated by the PI);
- The prospective subject or LAR will be provided with information and materials in a location appropriate to the study and offering the privacy necessary to ask questions about the study before deciding to participate;
- In obtaining informed consent, the PI will give the subject (or LAR) sufficient opportunity, commensurate with the risk level of the research, to consider whether or not to participate. Time should be allowed for questions and full discussion. Information about the study should be presented in a neutral, non-coercive manner and in a language readily understandable to the subject or LAR;
- Except as otherwise approved by the IRBMED, informed consent shall be documented by the use of a written consent form approved by the IRBMED and signed by the subject or the subject’s LAR. A copy shall be given to the person signing the form;
- The informed consent document used by the researchers must be the most recent version approved by IRBMED and is valid only after its approval by the convened board or through expedited review.

Assent of Children

Refer to IRBMED SOP Part 7.IV.C

Federal regulations (45 CFR 46.402 and 21 CFR 50.3(n)) require that IRBMED determine that adequate provisions are made for soliciting the assent of children involved as study subjects when, in the judgment of the IRB, the children are capable of providing assent (i.e., a child’s affirmative agreement to participate in research). Mere failure to object will not, absent affirmative agreement, be construed as assent. The assent process will determine when children are capable of assent based on age and maturity of the children, psychological state of the children, and nature of the proposed research activity.

(2) Short Form ICD

Refer to HRPP OM Part 3.III.C.6.e.2
A short form written ICD may be used in certain circumstances (45 CFR 46.117(b)(2)). The short form consent process requires that the elements of informed consent required by HHS and/or FDA regulations are presented orally to the subject or the subject’s LAR in the presence of a witness. For subjects who do not speak English, the witness must be conversant in both English and the language of the subject.

The IRBMED must approve the short form and a written summary (oral script) of what will be said to the subject or subject’s LAR.

The short form must include signature lines for the subject, or subject’s LAR and the witness.

The subject or subject’s LAR, the researcher consenting the subject, and a translator, if appropriate must sign the short form and/or the oral script according to the following table:

<table>
<thead>
<tr>
<th>Forms Required to Sign</th>
<th>Subject</th>
<th>Translator (when needed)</th>
<th>Researcher</th>
<th>Witness</th>
</tr>
</thead>
<tbody>
<tr>
<td>Forms</td>
<td>Short form</td>
<td>Recommended but not required UNLESS also serving as the witness or the researcher</td>
<td>Oral script</td>
<td>Short form and oral script</td>
</tr>
<tr>
<td>Required to Receive or Keep</td>
<td>A copy of the signed short form and oral script</td>
<td>Nothing UNLESS also serving as the researcher</td>
<td>Original signed short form and signed oral script</td>
<td>Nothing</td>
</tr>
</tbody>
</table>

The subject or subject’s LAR will receive a copy of the signed short form and oral script.

Refer to additional guidance on the IRBMED website.

(3) Informed Consent Waivers, Alterations, Exceptions and Substitutions

Waiver or Alteration of Informed Consent

The IRBMED may approve a consent procedure which does not include or which alters some or all of the basic elements of informed consent or which waives the requirement to obtain informed consent, if the IRBMED finds that appropriate conditions of 45 CFR 46.116(c) or (d) are satisfied and the research is not under FDA oversight.

Waiver of informed consent may be approved by the IRBMED only if it finds and documents that:

- The research involves no more than minimal risk to subjects;
- The waiver or alteration will not adversely affect the rights and welfare of the subjects;
- The research could not practicably (i.e. feasibly) be carried out without the
waiver of alteration; and

• Whenever appropriate the subjects will be proved with additional pertinent information after participation.

Alternatively, the IRB may approve a waiver or alteration of consent if it determines and documents that:

• The project is to be conducted by or subject to the approval of state or local government officials; and

• The project is designed to study, evaluate, or otherwise examine:
  o Public benefit or service programs;
  o Procedures for obtaining benefits or services under those programs;
  o Possible changes or alternatives to those programs or procedures; or
  o Possible changes in methods or levels of payment for benefits or services under those programs; and

• The research could not practicably be carried out without the waiver or alteration.

Projects involving the use of deception in the consent process must meet the same criteria as required for waiver of informed consent.

Waiver of Requirement for Parental Permission

For research involving children as human research subjects, IRBMED may waive the requirement to obtain parental permission if it determines and documents that:

• The research involves no more than minimal risk to the subjects;

• The waiver or alteration does not adversely affect the rights and welfare of the subject;

• The research cannot practicably be carried out with the waiver or alteration;

• When appropriate, the subjects will be provided with additional pertinent information after participation; and

• The research is not FDA-regulated.

Alternatively, the IRBMED may waive the requirement to obtain parental permission if it determines and documents that:

• The research is designated for conditions or for a subject population for which parental or guardian permission is not a reasonable requirement to protect the subjects;

• An appropriate mechanism for protecting the children who will participate as subjects is substituted; and

• The research is not FDA regulated.

Waivers of Documentation of Informed Consent

“Waiver of documentation” is a regulatory term that means the informed consent process takes place but the requirement to “document” that process does not involve obtaining subjects’ signatures on a written document. The IRBMED may waive the requirement for the PI to obtain a signed ICD for some or all of the
subjects if the requirements of 45 CFR 46.117(c) and/or 21 CFR 56.109(c)(1) are satisfied:

- The only record linking the subject and the research would be the ICD and the principal risk would be potential harm resulting from a breach of confidentiality (in which case each subject must be offered the opportunity to receive the documentation); or
- The research presents no more than minimal risk of harm to subjects and involves no procedures for which written consent is normally required outside of the research context.

Situations in which a waiver of documentation of informed consent is allowed include, but are not limited to:

- Telephone or web-based surveys
- Blood draws or urine collection (where HIPAA does not apply or can be waived);
- Research involving deviant or illegal behavior;
- Research involving socially sensitive issues, such as HIV status.

When the IRBMED waives the requirement for documentation of informed consent, the required elements of informed consent must be conveyed to the subject verbally or by electronic or printed text. Even though subjects do not sign a document, the IRBMED may still require that subjects be provided with written information about the study. The text of the written or oral informed consent scripts and any informational documents provided to subjects must be reviewed and approved by the IRBMED before their use.

Emergency Research Exception from Informed Consent (EFIC)

Refer to HRPP OM Part 3.III.C.6.e.iii.

Refer to IRBMED Guidance – Emergency Research with Exception From Informed Consent on the IRBMED website.

Refer to IRBMED Position Statement January 11, 2007 on the IRBMED website.

Emergency Research with Exception from Informed Consent (EFIC) is planned research conducted where subjects are in an emergent need of clinical care. “Emergency Research” is distinct from “Emergency Use Research”, in that the latter is an unplanned need to use an investigational agent that arises emergently for a single patient/subject.

In the course of its review, approval, and continuing review of clinical research proposals, the IRBMED may approve a research proposal without requiring that informed consent of all research subjects is obtained prior to the commencement of the research. The IRBMED (with the concurrence of a licensed physician who is a member of, or consultant to, the IRBMED and who is not otherwise participating in the clinical investigation) must find and document under 21 CFR 50.24 for an EFIC project each of the following:

(i) The human subjects are in a life-threatening situation, available treatments are unproven or unsatisfactory, and the collection of valid scientific evidence, which may include evidence obtained through randomized placebo-controlled investigations, is necessary to determine the safety and effectiveness of particular interventions.
(ii) Obtaining informed consent is not feasible because:

- The subjects will not be able to give their informed consent as a result of their medical condition;
- The intervention under investigation must be administered before consent from the subjects' LAR is feasible; and
- There is no reasonable way to prospectively identify the individuals likely to become eligible for participation in the clinical investigation.

(iii) Participation in the research holds out the prospect of direct benefit to the subjects because:

- Subjects are facing a life-threatening situation that necessitates intervention;
- Appropriate animal and other preclinical studies have been conducted, and the information derived from those studies and related evidence support the potential for the intervention to provide a direct benefit to the individual subjects; and
- Risks associated with the investigation are reasonable in relation to what is known about the medical condition of the potential class of subjects, the risks and benefits of standard therapy, if any, and what is known about the risks and benefits of the proposed intervention or activity.

(iv) The clinical investigation could not practicably be carried out without the exception from informed consent requirements.

(v) The proposed investigational plan defines the length of the potential therapeutic window based on scientific evidence, and the Principal Investigator has committed to attempting to contact a LAR for each subject within that window of time and, if feasible, to asking the LAR contacted for consent within that window rather than proceeding without consent. The Principal Investigator summarizes efforts made to contact the LAR and makes this information available to the IRBMED within fourteen (14) business days of each subject enrollment without consent.

(vi) The IRBMED reviews and approves informed consent procedures and an acceptable informed consent document. These procedures and the informed consent document are to be used with subjects or their LAR in situations where use of such procedures and documents is feasible. The IRBMED reviews and approves procedures and information to be used when providing an opportunity for a family member to object to a subject's participation in the clinical investigation.

(vii) Additional protections of the rights and welfare of the subjects will be provided, including, at least:

- Consultation (including, where appropriate, consultation carried out by the IRBMED) with representatives of the communities in which the
clinical investigation will be conducted and from which the subjects will be drawn;

- Public disclosure to the communities in which the clinical investigation will be conducted and from which the subjects will be drawn, prior to initiation of the clinical investigation, of plans for the investigation and its risks and expected benefits;

- Public disclosure of sufficient information following completion of the clinical investigation to apprise the community and researchers of the study (including the demographic characteristics of the research population) and its results;

- Establishment of an independent data monitoring committee to exercise oversight of the clinical investigation; and

- If obtaining informed consent is not feasible and a LAR is not reasonably available, the Principal Investigator has committed, if feasible, to attempting to contact within the therapeutic window the subject’s family member who is not a LAR, and asking whether he or she objects to the subject's participation in the clinical investigation. The Principal Investigator will summarize efforts made to contact family members and make this information available to the IRBMED within fourteen (14) days of subject enrollment without consent.

(4) Research Subject to FDA Regulations – Waiver or Alteration of Informed Consent

Refer to HRPP OM Part 3.III.C.6.e.iv.

(5) Research Subject to Both HHS and FDA Regulations

Refer to HRPP OM Part 3.III.C.6.e.v.

(6) Research Subject to HIPAA Regulations

Waiver of HIPAA Authorization

Under HIPAA, researchers must obtain written authorization from a research subject for release of protected health information that the researcher will collect, use or disclose for the study.

A PI may submit for review and approval a waiver of HIPAA authorization for the use and disclosure of PHI for research purposes, which may include a database or registry, or other human subject research projects not otherwise subject to IRBMED oversight, in which PHI may be used or disclosed without patient authorization, including decedents, PHI and review of PHI preparatory to research (e.g., Certification Preparatory to Research)

Depending on the type of submission (e.g., independent of or as part of an initial application in eResearch), the Privacy Board or full convened IRBMED Board may approve a waiver only if both of the following criteria are met:

- The project has been submitted to the IRBMED for review and has been deemed to be exempt from ongoing IRBMED oversight; and
- The use or disclosure of PHI for the project involves no more than minimal
risk of harm to the privacy of individuals. This criterion may be met where all of the following elements are present:

- An adequate plan is in place to protect patient identifiers and PHI from improper use and disclosure; and
- An adequate plan is in place to destroy the identifiers at the earliest opportunity consistent with the conduct of the research, unless there is a Privacy Board-approved health or research justification for retaining the identifiers or such retention is otherwise required by law; and
- Adequate written assurances that the PHI will not be reused or disclosed to any other person or entity, except as required by law, for authorized oversight of the research study, or for other research for which the use or disclosure would be permitted by HIPAA; and
- The Waiver of Authorization will not adversely affect the rights and welfare of the subjects; and
- The research could not practicably be conducted without the Waiver of Authorization; and
- The research could not practicably be conducted without access to and use of the PHI; and
- Whenever appropriate, the subjects (including their physicians, as applicable) are provided with additional pertinent information after participation; and
- Where the Principal Investigator anticipates the disclosure of PHI outside the Covered Entity (as that may be determined from time to time), the Principal Investigator must account for each disclosure and retain records of such disclosures.

Privacy Board

The Privacy Board operates under the authority of and in accordance with HIPAA (45CFR164) and applicable University policies and procedures. The Privacy Board is authorized to review and approve, if found to be in compliance with the requirements of these SOPs and HIPAA, the following:

- Waivers of authorization for research not subject to the Common Rule, or exempt from IRBMED oversight under the Common Rule, including waivers in connection with development and maintenance of research databases and registries;
- Investigator’s application for Certification Preparatory to Research for reviews of protected health information;
- Information (PHI) preparatory to research submitted in the eResearch application (also known as “pre-review of clinical data sets” in the initial application in eResearch);
- Investigator’s application for Certification Preparatory to Research for research involving decedents’ PHI submitted in the eResearch application;
- In consultation with other units (e.g., the UMHS Privacy Office and DRDA), any use or disclosure of limited data sets under data use agreements; and
- Waivers of Authorization for research may also be granted by the IRBMED Board as part of a review for a project under IRBMED jurisdiction.

Privacy Board Membership

Privacy Review Board Membership includes, but is not limited to, the following members:
• **Chair** – The Privacy Board Chair is an experienced and active member of the IRBMED and is well-informed concerning the laws, regulations, and University policies and procedures that govern the use or disclosure of protected health information in connection with human subjects’ research.

• **Voting member(s)** – The Privacy Board shall include at least one member who is a “Community Representative”, who is not affiliated with UM or any entity conducting or sponsoring the research, and who is not a family member (e.g., spouse, domestic partner or dependent) to any person affiliated with the University or an entity conducting the research.

**Procedures for Waiver of Patient Authorization**

All applications for Waiver of Authorization require review by the Privacy Board or by the convened IRBMED board where the request is part of an initial application, unless eligible for expedited review. All other applications will be reviewed and approved by the IRBMED Director or designated Privacy Board Administrator, unless consultation with other units (i.e., the UMHS Privacy Office or ORSP) is necessary.

Upon receipt of an application from a PI requiring Privacy Board review the regulatory staff will forward a complete set of documents (in electronic and/or printed form) directly relevant to the application to the Privacy Board members. All of the project documents shall be made available to Privacy Board members and authorized consultants for advance review.

Prior to the Privacy Board meeting, the regulatory staff may request from the PI additional information or documents, or explore proposed revisions that may be required as a condition of Privacy Board approval of the project.

The designated regulatory staff member will present the project to the Privacy Board membership at a convened meeting, unless approval is expeditable as provided below.

An application for Waiver of Authorization for Use or Disclosure of PHI in connection with a research project may be approved by expedited review if the research involves no more than minimal risk of violation of the privacy of the individuals who are the subject of the research for which use or disclosure is being sought. Expedited review may be carried out by the Chair or any member of the Privacy Board as designated by the Chair.

A reviewer has the authority to approve an expeditable project without a vote of the convened Privacy Board. If the reviewer believes that the application is not suitable for expedited review, the reviewer shall defer any decision and instead submit the application to a full review by the Privacy Board.

All applications approved by expedited review are available to the Privacy Board Members, the UMHS Privacy Office, and authorized consultants on an ongoing basis via eResearch.

Refer to [HRPP OM Part 3.III.C.6.g](#) for a detailed description of points the IRBMED should consider in determining whether a protocol includes plans sufficient to address privacy and confidentiality concerns.

**f) Data Monitoring**

Refer to [HRPP OM Part 7.II.](#)

Refer to IRBMED website - [Data Safety Monitoring Boards: Agency for Healthcare](#)
Research and Quality (AHRQ), Mandatory Use Guidance

With respect to any research project or class of research projects, the IRBMED may impose additional conditions on the conduct of the research at any time prior to, concurrent with, or following approval, when in the judgment of the IRBMED such additional conditions are necessary or appropriate for the protection of human research subjects.

(1) Considerations for the Imposition of Special Monitoring Requirements

The IRBMED may, at its discretion, perform monitoring or request monitoring from the Office of Human Research Compliance Review (OHRCR) – via UMOR – of a project; in addition to information received through the initial application, any amendments, annual continuing reviews, and analyses of interim reports, such as adverse events and audit reports. For example, the IRBMED may choose to undertake extra monitoring for research which presents greater than minimal risk, to gauge the progress of recruitment for vulnerable subjects, or to follow the research progress on controversial subject matter. The IRBMED may also consider the frequency and nature of AEs reported to date.

The IRBMED may also choose to monitor one or more of the projects of a single Principal Investigator in consideration of the experience of the Principal Investigator or as follow-up to previous reports of complaints, non-compliance, or prior IRBMED interactions with the individual.

(2) Examples of Special Monitoring Requirements

Monitoring may include, but is not limited to requesting copies of access to any or all of the following from the PI:

• Signed informed consent documents;
• Site visits to research locations;
• Shortened approval periods and/or interim scheduled reports from the Principal Investigator during the approval period;
• Interviews with subjects;
• Interviews with study personnel;
• Drug dispensing/IDS logs;
• Third party witness to the informed consent process;
• Study files and research records;
• Independent Data and Safety Monitoring Board (DSMB) reports; and
• Other independent third-party monitoring reports

The IRBMED shall communicate with the PI as appropriate, regarding the outcomes of these additional monitoring efforts.

g) Privacy and Confidentiality Protection

Refer to HRPP OM Part 3.III.C.6.g.


Refer to University of Michigan Website Guidance for Sensitive Human Subjects Data: http://safecomputing.umich.edu
To approve research, the IRBMED must determine that, where appropriate, there are adequate provisions to protect the privacy of subjects and the confidentiality of data. Regulatory, institutional, and IRB policies and guidance are used to confirm that the protocol appropriately and adequately protects privacy.

The PI must include a plan to protect participants’ privacy and confidentiality in the eResearch application, protocol or other documents submitted to the IRB to include a description of the types of privacy and confidentiality information that the researcher must include in its plan. The IRB reviews the PI’s plan to protect participants’ privacy and confidentiality to assess the adequacy of the protection.

In reviewing confidentiality protections, the IRBMED shall consider the nature, probability, and magnitude of harms that would be likely to result from a disclosure of collected information outside the research context. It shall evaluate the effectiveness of proposed techniques to anonymize, code, encrypt, store, or access the information, and any other relevant factor in determining the adequacy of confidentiality protections.

h) Vulnerable Subjects

Refer to HRPP OM Part 7.IV.
Refer to IRBMED SOP Part 7.IV

Special federal regulations apply to research involving vulnerable populations. These groups include, but are not limited to:

- Children (individuals who have not attained the legal age for consent to the treatments or procedures involved in the research, under the applicable law of the jurisdiction in which the research will be conducted (45 CFR 46 Subpart D; 21 CFR 50 Subpart D);
- Pregnant women, fetuses, and neonates, including those of uncertain viability (45 CFR 46 Subpart B);
- Prisoners:
  - Individuals involuntarily confined or detained in a penal institution, including:
    - Individuals detained in other facilities by virtue of statutes or commitment procedures which provide alternatives to criminal prosecution or incarceration in a penal institution, as well as individuals detained pending arraignment, trial, or sentencing (45 CFR 46 Subpart C).
  - Individuals who are cognitively impaired or lack decision-making capacity (21 CFR 50.20);
- Individuals who otherwise may be subject to coercion or undue influence (e.g., economically or educationally disadvantaged persons, employees or students of investigators conducting the study, or patients of physician-investigators (21 CFR 50.20).

When members of any of these groups participate in research, the IRBMED requires investigators to specify what additional protections, if any, will be provided to these persons to protect their rights and welfare (e.g., to minimize risks unique to these groups and the possibility of coercion or undue influence). In reviewing these research projects, the IRBMED will ascertain that inclusion of a vulnerable population
is adequately justified and that safeguards are implemented to minimize risks unique to that population.

Laws governing research involving vulnerable populations, including laws on who may consent on behalf of children or cognitively impaired or incapacitated adults, vary from state to state.

**Michigan Law**

Michigan Law requires special consent for procedures or treatments for the following conditions:

- breast cancer
- electroconvulsive therapy
- HIV/AIDS testing
- pregnancy termination
- surgery for mental health patients
- terminal illness

*Refer to IRBMED SOP Part 11* for references to guidance for determining the nature of requirements applicable to Michigan and to research proceeding outside of Michigan.

*Refer to HRPP OM Part 11.II* for a detailed description on Michigan requirements and guidance for determining the nature of requirements applicable to research proceeding outside of Michigan.

i) **Test Article Accountability Procedures**

*Refer to HRPP OM Part 8.VIII.D*

*Refer to HRPP OM Part 3.III.C.6.i*

j) **Resources**

*Refer to HRPP OM Part 3.III.C.6.j*

IRBs will determine that research studies have the resources necessary to protect subjects by evaluating all of the following:

- There is adequate time for the investigators to conduct and complete the research;
- There are an adequate number of qualified staff;
- The facilities where the research will be conducted are adequate;
- PIs have access to a population that will allow recruitment of the necessary number of subjects; and
- Medical or psychosocial resources that subjects may need as a consequence of the research are available.

7. **IRBMED Review and Monitoring of FDA-Regulated Research**

*Refer to HRPP OM Part 8*

*Refer to HRPP OM Part 3.III.C.7.*

*Refer to IRBMED SOP Part 8*
D. IRB Administrative Functions

1. IRBMED Meetings

a) An IRB must review proposed research and conduct continuing reviews at convened meetings at which a majority of the members of the IRB are present.

b) At convened meetings at least one non-scientist member must be present in order to meet quorum; at least one unaffiliated member, who represents the general perspective of subjects, should be present at the majority of meetings in a given year. Attendance of all present members is recorded in the meeting minutes.

c) In order for the research to be approved, it must receive approval by majority vote of the quorum (as described above). If, during the course of the meeting, quorum is lost, votes may not be taken until it has been restored.

d) When convened-board review is not required, the SOPs must include details of any process, such as expedited review procedures (as described above) or subcommittee procedures, which may be used to supplement the IRB's review responsibilities.

e) IRB members may agree, during an appropriately convened meeting, to issue conditional approval for a project only if any requested clarifications or modifications are not relevant to the determinations required by the IRB under the Common Rule or its Subparts (45 CFR 46) or, as applicable, FDA regulations (21 CFR 56). If substantive clarifications or modifications regarding the protocol or informed consent documents are required as a condition of approval, approval must be deferred pending subsequent review of responsive material by the convened IRB of responsive material.

f) Although it is preferred that IRB meetings take place with all participating members physically present, an IRB may establish protocols to convene meetings via video conference, teleconference or similar means. Such protocols must provide a means for all participants to receive the meeting materials prior to the meeting and facilitate active and equal participation in the discussion of all protocols. The protocols must further provide that minutes from meetings convened in this manner will reflect that these two conditions have been met, in addition to other mandated information (e.g., presence of a quorum including at least one non-scientist, discussion and resolution of problematic issues, final vote).

2. Notification of Decisions

a) An IRB will notify investigators in writing of its decision to approve or disapprove a proposed research activity or of modifications to the proposal that are required to secure IRB approval.

b) If the IRB decides to disapprove a research activity, it must include a statement of the reasons for its decision in its written notification and must give the investigator an opportunity to respond in person or in writing.

c) An IRB will notify the IO or DIO and other institutional officials, when appropriate, of its decisions regarding proposed research activities by formal or informal means, such as through access to relevant electronic databases.
3. IRB Response to Noncompliance, ORIOs and Other Required Reporting

Refer to IRBMED SOP Part 12

4. IRB Records and Reports

Reports Uploaded into eResearch

The following documentation associated with eResearch submissions, determinations and tracking will be either uploaded (and permanently archived) to the appropriate submission itself, or via AE (Adverse Event Report) or ORIO (Other Reportable Information or Occurrences) Reports:

- AEs are reported according to the AE Reporting guidance available on the IRBMED Website;
- Statements of significant new findings provided to subjects;
- Documentation of exemption determinations, include the category by which research was determined to be exempt;
- Documentation of approvals using the expedited procedure, including the applicable criteria by which the research was approved using the expedited procedure;
- Description of action taken by a reviewer;
- Records of continuing review activities
- Progress reports submitted by PIs;
- Submission approval letters;
- Documentation of Institutional Authorization Agreements, Individual Investigator Agreements, or Collaborating Institutional Agreements
- Minutes of IRBMED meetings sufficiently detailed to show:
  1. Attendance at meetings;
  2. Actions taken by the IRB;
  3. The vote on these actions including the number of members voting for, against, and abstaining;
  4. The basis for requiring changes in or disapproving research;
  5. A written summary of the discussion of problematic issues and their resolution;
  6. Separate deliberations for each action;
  7. When an alternate member replaces a primary member;
  8. For initial and continuing review, the approval period;
  9. The names of IRB members who leave the meeting because of a conflict of interest along with the fact that a conflict of interest is the reason for the absence;
  10. Unless documented in the IRB records, determinations required by the regulations and protocol-specific findings justifying those determinations for:
      a. Waiver or alternation of the consent process,
      b. Research involving pregnant women, fetuses, and neonates,
      c. Research involving prisoners, and
      d. Research involving children;
11. When following DHHS regulations or guidance, documentation of justification of any deletion or substantive modification of information concerning risks or alternative procedures contained in a DHHS-approved sample consent document; and

12. When following FDA regulations or guidance, documentation of the rationale for significant risk/non-risk device determinations.

Reports and Communications Archived on Network

- A membership roster of the current IRBMED members (and alternates) and their qualifications (degrees earned, area of expertise, membership role.) sufficient to describe each member's anticipated contribution to IRBMED deliberations and any employment relationship between members and UM.
- Written SOPs
- Documentation of member and staff training.

Retention

Refer to [IRBMED Guidance – Record Retention](#) on the IRBMED website for information on the length of time study records are to be kept on file.

Hard copy materials are logged and stored off-site in a secure manner in a commercial storage facility. Retrieval of documents stored off-site is arranged by contacting the IRBMED office, who will notify the facility for the appropriate records to be delivered on an assigned date.

- IRBs must maintain records for at least three (3) years after the completion of a research study.
- If a research application is terminated without subject enrollment, IRB records must be maintained for at least three (3) years following termination;
- If an IRB performs functions on behalf of a "covered entity (such as the University of Michigan Hospitals and Health Centers) related to HIPAA and research, those records must be retained for at least six (6) years, either by the IRB, or by the covered entity; and
- Administrative units responsible for IRB operations may impose longer retention and specific destruction standards.

Inspection of Records

Paper and electronic documents will be made accessible for inspection and copying by authorized representatives of UM, relevant sponsors, and government authorities with jurisdiction (such as OHRP, FDA, and NIH) at reasonable times and in a reasonable manner.

E. Quality Assurance and Quality Improvement

Refer to [HRPP OM Part 12.I.A-B](#)

SOPs

The IRBMED cooperates with the Medical School Associate Deans for Research and Regulatory Affairs, UM Health System (UMHS) Compliance Committee, and the OVPR to establish, review, and revise these SOPs. These SOPs and any substantive revisions thereto, are subject to review and approval by the Medical School Associate Deans for Research and Regulatory Affairs, UMHS Compliance Committee, and the VPR. Non-
substantive revisions such as modifications to enhance regulatory flexibility and workflows, inclusion of standard forms, guidance documents, and similar information developed by the IRBMED in consultation with the Medical School Associate Deans for Research and Regulatory Affairs and OVPR do not require further review or approval. Outdated sections of these SOPs will be archived in such a way that changes and dates of approval may be followed.

The IRBMED SOPs will be reviewed at least once every five (5) years in conjunction with the AAHRP accreditation cycle. UMOR initiates a comprehensive review of the HRPP Operations Manual at the same time. IRB SOPs must make provisions for such a review of SOPs on the same cycle, or more frequently, at the IRB’s discretion.

Revisions to SOPs may be made at any time, as required by changes in law, ethical standards, institutional policy, quality assurance activities or other considerations.

Substantive revisions require advance approval by the HRPP Director.

IRBMED SOPs are conducted by the Medical School Associate Deans for Research and Regulatory Affairs, UMHS Compliance Committee, and UMOR.

Internal Quality Assurance
The IRBMED routinely conducts internal review of its staff and board member operations, as well as reviewing the human subject research application (eResearch) operations, as part of its continuous quality improvement efforts, in order to measure the effectiveness of its human research protection program and to determine if its review processes are performed and recorded in compliance with established standards.

Review will be conducted periodically by the following means:
- Solicitations in writing or by survey of the IRBMED Chairs, members, staff, and affiliated PIs and study team personnel as well as from standing and ad hoc research advisory councils within the jurisdiction of the IRBMED;
- Consultation services by the Chairs, members, or staff, followed by requests for feedback regarding these consultative meetings;
- Peer assessment;
- Periodic checks for quality improvement;
- Review by other institutional units, such as OHRCR.

IV. OTHER REVIEW UNIT STANDARD OPERATING POLICIES AND PROCEDURES

Refer to HRPP OM Part 3.IV