

SOP: Minutes

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1. PURPOSE

- 1.1. This procedure establishes the process to take IRB minutes.
- 1.2. This procedure begins when the meeting is called to order.
- 1.3. This procedure ends when the minutes are finalized.

2. POLICY

- 2.1. None

3. RESPONSIBILITY

- 3.1. HRPP staff members carry out these procedures.

4. PROCEDURE

- 4.1. Use the “Minutes (HRP-510)” template to record minutes.
- 4.2. Record at the beginning of the minutes:
 - 4.2.1. “Members Present”: Record the following information on IRB members present at any time during the meeting and having voting status at least once during the meeting¹:
 - 4.2.1.1. Name.
 - 4.2.1.2. Status²
 - 4.2.1.3. Whether the IRB member is an alternate
 - 4.2.1.4. Whether the IRB member attended by teleconference.
 - 4.2.2. “Others Present”: Record the following information on individuals present at any time during the meeting who never have voting status:³
 - 4.2.2.1. Name.
 - 4.2.2.2. Role
- 4.3. Record the total number of regular members on the current IRB roster and the number of members required for quorum⁴.
- 4.4. If IRB members are present by teleconference, indicate whether they received all pertinent material before the meeting and were able to actively and equally participate in all discussions
- 4.5. Record the time the meeting is called to order.
- 4.6. Record a summary of the discussion of items unrelated to the review of specific research.
- 4.7. For each item related to specific research:
 - 4.7.1. Record the type of review⁵
 - 4.7.2. Record relevant information about the research:
 - 4.7.2.1. Title
 - 4.7.2.2. Principal investigator
 - 4.7.2.3. IRB number

¹ If an IRB member has non-voting status for the entire meeting, list as an “Others Present.”

² For example: IRB chair, IRB vice-chair, scientific member, non-scientific member, unaffiliated member, pediatric experience, prisoner representative

³ This may include IRB members who are present for the meeting but never vote, consultants, non-IRB members, HRPP staff, etc.

⁴ The whole number greater than one-half of the number of regular members

⁵ For example: Initial, continuing, modification, <Unanticipated Problem Involving Risks to Subjects or Others>, <Serious Noncompliance>, <Continuing Noncompliance>, <Suspension of IRB Approval>, <Termination of IRB Approval>, study, site

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- 4.7.2.4. IND or IDE number, if any
- 4.7.2.5. HHS grant title and ID, if any
- 4.7.2.6. Documents reviewed
- 4.7.3. When needed for clarity, summarize previous IRB actions.
- 4.7.4. If any item is not acted upon, record the reason⁶.
- 4.7.5. If a consultant provided an oral report, summarize the key information provided.
- 4.7.6. If there were any controverted issues (IRB members expressed a difference of opinion), summarize the issue, label as a controverted issue, and summarize the resolution, if any.
 - 4.7.6.1. If there were no controverted issues, record this.
- 4.7.7. Record the motion.
 - 4.7.7.1. For a motion of "Approve" or "Conditionally Approve" related to an initial or continuing review submission record:
 - 4.7.7.1.1. The approval period or that continuing review is not required.
 - 4.7.7.1.1.1. If continuing review is not required by "WORKSHEET: Criteria for Approval (HRP-400)" but the IRB requires continuing review, document the rationale for requiring continuing review.
 - 4.7.7.1.2. Whether the risk is <Minimal Risk> or greater than <Minimal Risk>
 - 4.7.7.1.3. Any required checklist determinations along with study-specific findings supporting those determinations
 - 4.7.7.1.4. Any rationale for any <Non-significant Risk Device> or <Significant Risk Device> determination
 - 4.7.7.1.5. Document that the IRB determined that the proposed research met the criteria for approval.
 - 4.7.7.1.5.1. In the case of a financial interest that is <Related to the Research> document instead that the IRB determined that proposed research with the management plan for the financial interest met the criteria for approval.
 - 4.7.7.2. For a motion of "Conditionally Approve" record the IRB's modifications required to secure approval and the reasons for those modifications.
 - 4.7.7.2.1. Document that the IRB determined that the proposed research with the requested modifications met the criteria for approval.
 - 4.7.7.2.1.1. In the case of a financial interest that is <Related to the Research> document instead that the IRB determined that proposed research with the requested

⁶ For example: Loss of all non-scientific members, missing expertise, meeting ended early due to fire alarm

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modifications and with the management plan for the financial interest met the criteria for approval.

- 4.7.7.3. For a motion of “Defer” record the IRB’s reasons and recommendations.
- 4.7.7.4. For a motion of “Disapprove” record the IRB’s reasons.
- 4.7.7.5. For a motion of “Suspend” record the specific activities suspended and the IRB’s recommendations, if any.
- 4.7.7.6. For a motion of “Lift Suspension” no other information needs to be recorded.
- 4.7.7.7. For a motion of “Terminate” record the IRB’s reasons.
- 4.7.8. Record the vote as the numbers:
 - 4.7.8.1. “For”: Voting for the motion.
 - 4.7.8.2. “Against”: Voting against the motion
 - 4.7.8.3. “Abstain”: Present for the vote, but not voting “For” or “Against”
 - 4.7.8.4. “Absent”: Not present for reasons other than a <Conflicting Interest>
 - 4.7.8.4.1. Record the names of absent members (members in attendance at the meeting, but absent from the room for the vote)
 - 4.7.8.5. “Recused”: Not present for discussion and voting due to a <Conflicting Interest>
 - 4.7.8.5.1. Record the names of recused members
 - 4.7.8.6. Non-Voting Status: Present at the meeting but not in voting status (in voting status for some items but not in voting status for all items)
 - 4.7.8.6.1. Record the names of members present in non-voting status
- 4.8. Record the time the meeting is adjourned.
- 4.9. Provide the minutes to the <Meeting Chair> for review and approval, and provide to the IRB as an information item.
- 4.10. Provide approved minutes to the [Organizational Official] and the IRB members who attended the meeting.
- 4.11. Upon request Florida Hospital makes IRB records (including minutes) available to clients provided they are relevant to the client. Such records may be excerpted and/or redacted to comply with Florida Hospital’s obligations to maintain confidentiality.

5. REFERENCES

- 5.1. 21 CFR §56.115(a)(2)
- 5.2. 45 CFR §46.115(a)(2)