# IRB SOP 302 IRB Materials for Review

# **Purpose**

This Standard Operating Procedure (SOP) describes the materials provided to the IRB for review purposes.

# **Policy**

IRB review materials are provided to IRB members seven calendar days in advance of convened meetings, except in special circumstances described in *SOP 301: IRB Meeting Preparation*. IRB members may request additional information or supporting documents at any time.

### **Procedures**

- 1.0 IRB Review Procedures
  - 1.1 Primary/Secondary Reviewers

The primary and secondary reviewer (if applicable) for a given research

#### 1.2 Other reviewers

All IRB members receive the IRB agenda, previous month's minutes for approval, appropriate IRB application(s), informed consent (or request to waiver informed consent) and surveys/questionnaires. Relevant materials are to be provided for all types of IRB review including initial review, continuing review and amendments for review at the convened meeting.

#### 2.0 Materials Provided to IRB Members for Review

#### 2.1 General

- 2.1.1 Meeting Agenda
- 2.1.2 Minutes for previous meetings
- 2.1.3 Report of completed expedited reviews
- 2.1.4 Educational materials

### 2.2 Initial Applications – Materials provided by the investigator (as applicable)

- 2.2.1 Application form(s)
- 2.2.2 Application supplements
- 2.2.3 Consent/assent
- 2.2.4 Recruiting materials
- 2.2.5 Data collection instruments
- 2.2.6 Investigator's drug brochure/package insert
- 2.2.7 Device brochure/other device information
- 2.2.8 Industry research: Protocol
- 2.2.9 Relevant grant applications/contracts
- 2.2.10 Financial Conflict of Interest disclosure/management plan
- 2.2.11 Other materials relevant to study or deemed useful by the IRB

# 2.3 Continuing Reviews

- 2.3.1 Continuing Review form
- 2.3.2 Updates to IRB Application Part A
- 2.3.3 Consent/assent documents
- 2.3.4 Relevant post-approval reports (e.g., Data Safety Monitoring reports)
- 2.3.5 Any other materials provided by the investigator
- 2.3.6 Other materials relevant to study or deemed useful by the IRB

#### 2.4 Amendments

- 2.4.1 Amendment Review form
- 2.4.2 Modified protocol
- 2.4.3 Modified consent/assent
- 2.4.4 Modified recruitment materials
- 2.4.5 Modified investigator brochure/package insert
- 2.4.6 Other modified study documents

# University Related Documents SOP 301: IRB Meeting Preparation