

Protocol Review & Monitoring Committee Review Types

If your Protocol is Cancer related
&
DMG approval obtained

FULL BOARD COMMITTEE REVIEW

Subject to a review by the board members at a scheduled meeting, with PI present, subject to meeting dates / deadlines; review of scientific merit, biostatistical plan analysis, and appropriate allocation of resources
(10 business days)

ADMINISTRATIVE REVIEW

Expedited review by an executive sub-committee as received. Not subject to meeting dates / deadlines
(5 business days)

All interventional studies in which individuals are assigned prospectively by an investigator based on a protocol to receive specific interventions. The participants may receive diagnostic, treatment, behavioral, or other types of interventions. The assignment of the intervention may or may not be random. The participants are followed and biomedical and/or health outcomes are assessed.

PRMC Approval

SUBMIT TO IRB

- National Cooperative group studies
- Externally Peer-Reviewed Trials
- Any peer-reviewed study supported by an NIH mechanism (ie: ROIs, UOIs, U10s, POIs, and P50s, etc.)
- Observational studies that focus on cancer patients and healthy populations and involve no prospective intervention or alteration in the status of the participants and assess biomedical and/or health outcome(s) in pre-defined groups of participants.
- Ancillary or Correlative studies
- Laboratory-based studies that are prospectively collecting consent and using specimens to assess cancer risk, clinical outcomes, response to therapies