



RESEARCH GOVERNANCE UNIT
St. Vincent's Hospital (Melbourne)
Caritas Christi Hospice
St. George's Health Service
Prague House
Cambridge House
DePaul House

MULTISITE APPROVAL

Statement of Intent and Outcomes

The St Vincent's Hospital Human Research Ethics Committee is committed to fulfilling Section 5 of The National Statement on Ethical Conduct in Human Research (2007 – Updated 2018) by reducing the duplication of ethical review where possible.

Definitions

ERM is defined as the Ethical Review Manager, which is the database utilized by the multisite approval process.

Reviewing HREC is defined as the HREC who is allocated to undertake an ethical review of the study.

Accepting HREC is defined as other HRECs who accept the ethical approval of the reviewing HREC without further ethical review.

Lead Site is defined as the single site responsible for the ethical submission of documents for a study.

Master Participant Information and Consent Form is defined as the universal template which must be used by all sites participating in the study. This document must not contain site specific information.

Site Specific Participant Information and Consent Form is the universal template which must be used by all sites participating in the study, which has been modified to include site specific information (i.e. institution name, investigators name, contact details)

Procedure

The Streamlined Ethical Review Process for multisite clinical trials is an initiative of the Victorian, New South Wales, Queensland, Western Australia, Tasmania and South Australian Governments. This allows clinical trials to be reviewed by one HREC with the decision mutually accepted by other certified centres. This ensures that ethical review is sought once only and that each participating site need only obtain governance authorisation prior to the commencement of the study via the submission of a Site Specific Assessment (SSA).

St Vincent's Hospital is committed to this multisite approval process, and is certified to both review and accept the approval of other centres for multicentre trials. At all times, participating sites must adhere to the Standard Operating Procedures which are available from: <https://www2.health.vic.gov.au/about/publications/policiesandguidelines/standard-principles-for-operation>

The accredited HRECs which participate in the multisite approval pathway are listed in the Memorandum of Understanding, which can be accessed from the Victorian Department of Health website: <https://www2.health.vic.gov.au/about/clinical-trials-and-research/health-and-medical-research/national-mutual-acceptance>

HREC Review

The lead site is responsible for submitting the ethics application to the reviewing HREC.

For St Vincent's Hospital (Melbourne) to provide ethical review, the following documents are required at the time of submission (electronic copies only):

Document	No of e-copies required
Cover Sheet	1
Fee Form for Ethical Review	1
Application Form: Human Research Ethics Application (HREA)	1
Victorian Specific Module	1
Participant Information Sheet and Consent Form (Master)	1
Investigators Brochure (or Product Information)	1
Clinical Protocol	1
Budget	1
Questionnaires, Patient Facing Documents etc	1
Radiation Reports	1
CV & GCP of Coordinating Principle Investigator (CPI)	1
CV's of all Associate Investigators (AI) / Site Principal Investigators (PI's)	1
CTN	1 CTN Draft
Indemnity - HREC review only	1 signed original
Insurance Certificate	1 signed original
Privacy Declaration for External Researchers (as applicable)	1 signed original

If St Vincent's Hospital is listed as a site, the following documents are also required for governance review:

Document	Number of e-copies required
Site Specific Assessment Form	1
Participant Information Sheet and Consent Form (Site Specific)	1
CV & GCP's of PI and AI's at SVHM	1
Clinical Trial Research Agreement / Research Collaboration Agreement (as applicable)	1
Indemnity - Standard	1 signed original

Once a valid submission is received by the reviewing HREC, the Principal Investigator will be formally notified in writing to confirm the validity of the study and to confirm the meeting date at which the study will be reviewed. ERM will also be updated to ensure the study details are entered, to confirm the validity of the study, and to allocate the study to a meeting. The clock must commence running at this stage.

Once ethical review has occurred, the Principal Investigator will be formally notified of the outcome in writing. ERM must also be updated with the outcome, and the clock stopped.

Once the study has been ethically approved, a formal letter of approval must be sent to the Principal Investigator, listing the approved documents. This will be sent electronically, to allow the Principal Investigator to circulate the documents to the Sponsor (as applicable) Investigators at other participating sites, and the Research Governance Offices at each site where the study will be conducted.

Once approval is granted, other sites should commence the submission of SSA applications to other participating sites, using the approved documents as templates.

SSA Review

Once ethical approval has been obtained from an accredited HREC, a Site Specific Assessment application must be submitted by each participating site to their respective Research Governance Office, for governance review only.

SSA Applications must include the letter of approval from the lead site, all ethically approved documents, a site specific PICF and any legal documents including Clinical Trial Agreements, Insurance Certificates and CTN forms.

Once governance authorization has been granted, the site Principal Investigator will be notified in writing. ERM must also be updated to record the authorization.

Amendments

Where St Vincent's Hospital is the reviewing HREC, a full submission of the amendment is required for review and formal ethical approval. Once the amendment has been approved, a formal letter of approval must be sent to the Principal Investigator, listing the approved documents. This must be sent electronically.

Where St Vincent's Hospital is the accepting HREC, a full submission of the amendment along with the HREC approval letter is required for governance review. Once the amendment has been approved, a formal letter of approval must be sent to the site Principal Investigator.

Adverse Events

Where St Vincent's Hospital is the reviewing HREC, a full submission of any related, potentially related or possibly related Suspected Unexpected Serious Adverse Reaction (SUSAR) as well as line listings and annual safety reports are required for review and formal ethical approval.

Once the adverse event has been approved, a formal letter must be sent to the Principal Investigator. This must be sent electronically.

All adverse events occurring at St Vincent's Hospital must be reported to the HREC as soon as possible.

These events must be reported to the Victorian Managed Insurance Authority (VMIA) by the Research Governance Unit to comply with local insurance requirements.

Where St Vincent's Hospital is the accepting HREC, only those adverse events occurring at Vincent's Hospital or those that have an impact on the conduct of the trial must be reported as soon as possible.

Associated Procedures/Instructions

Procedure 5.13 – Monitoring Approved Research

Procedure 5.7 – Document and Record Management

Procedure 5.11 – Minimising Duplication of Ethical Review

Reference Documents

- The National Statement on Ethical Conduct in Research Involving Humans in accordance with the NHMRC Act, 2007 – Updated 2018 (Cth)
- Australian Code for the Responsible Conduct of Research (2018)

Authorized by:



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