# Terms Of Reference

**Metro South Hospital and Health Service**  
**Human Research Ethics Committee EC00167**

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PURPOSE:
The Metro South Human Research Ethics Committee (HREC), (EC00167) within Metro South Hospital and Health Service, is constituted and functions in accordance with the National Health & Medical Research Council (NHMRC) “National Statement on Ethical Conduct in Human Research (2007, updated 2018)” and complies with the “Australian Code for Responsible Conduct of Research (2018), the Queensland Health (QH) Research Management Policy and Framework (QHRMP; 2015) and the Queensland Human Rights Act (2019).

The HREC, thereafter known as ‘the Committee’, acts in a consultative and advisory capacity with researchers to ensure that all clinical, research and management practices are conducted in an ethical and scientifically robust manner. Key objectives of the Committee are to:

- Safeguard the mental and physical welfare, rights, dignity and safety of participants involved in human research;
- Facilitate and promote high calibre ethical research through efficient and effective review processes; and
- Ensure that all clinical and health research is conducted responsibly and in the interests of the wider community.

1. Scope of Responsibilities and Functions (National Statement section 5.1.27)

The Metro South HREC is appointed by and acts in an advisory capacity to the Metro South Hospital and Health Service. The Committee considers all research protocols involving humans across the total breadth of health services provided at the Princess Alexandra Hospital, Logan and Beaudesert Hospitals, Redland Hospital and Queen Elizabeth II Jubilee Hospital, as well as Community and Primary, Oral and Mental Health Facilities within the Metro South Hospital and Health Service.

The Metro South Chief Executive or Delegate, upon recommendation of the Committee, may grant approval for research proposals conducted within its facilities. Furthermore, the Committee reviews multi-centre research from other Hospital and Health Services, Jurisdictions and practices (including private health facilities). The Committee has been certified by the NHMRC to conduct reviews of multicentre research conducted within jurisdictions participating in the national mutual acceptance model and participates in the Queensland Health single ethical review process. To this end the Metro South HREC aims to:

- Provide balanced, independent, and timely review of research protocols involving human participants in respect to their ethical acceptability and scientific merit.
- Review and recommend approval of research which may facilitate expedited HREC approval at other participating Hospital and Health Service/s subject to individual HREC site specific requirements and low risk expedited review procedures.
- Oversee approved protocols during the course of the research until completion to ensure that they comply with approved ethical standards, legislation, codes of practice and policies. Specific monitoring of the conduct of research will be conducted via the Research Governance Office (RGO) and in the case of multi centre research, the Co-ordinating Principal Investigator at the respective Institution.
The HREC may obtain expert opinion or establish sub-committees to provide scientific/technical assessment and safety evaluation of research protocols along with compliance with regulatory requirements.

The HREC administrative team will register all research protocols submitted to the Metro South HREC along with any monitoring and reporting requirements, and approval of protocol amendments during the course of the research.

2. Relationships and Reporting

The Metro South HREC, established by the Institution in keeping with the National Statement (2007, updated 2018 section 5.1.27) will:

- Report to the Metro South Hospital and Health Service (HHS) Chief Executive via the Chair of Metro South Research. Formal mechanisms of reporting include the HREC Annual Compliance Report to the National Health and Medical Research Council (NHMRC), minutes for all HREC meetings signed off by the Chairperson, routine meetings, and an annual report for the Metro South Research Report.

- Review and recommend approval of research undertaken within and in collaboration with the Metro South HHS along with multi-centre research to be conducted in other Hospital and Health Services, jurisdictions and practices.

- Review and recommend approval of research which may facilitate expedited HREC approval at other participating HHS, subject to individual site specific requirements.

- Oversee the monitoring of approved research until completion and the provision of final reports to ensure that the research has complied with approved ethical standards along with relevant legislation, regulations, codes of practice, policies and procedures.

3. HREC Composition and Appointment (National Statement sections 5.1.29 – 5.1.36)

The membership of the HREC is constituted according to the National Statement and includes the following:

(a) Chair and Deputy Chair, with suitable experience, whose other responsibilities will not impair the HREC’s capacity to carry out its obligations under this National Statement;

(b) at least two lay people, one man and one woman, who have no affiliation with the institution and do not currently engage in medical, scientific, legal or academic work;

(c) at least one person with knowledge of, and current experience in, the professional care, counselling or treatment of people; for example, a nurse or allied health professional;

(d) at least one person who performs a pastoral care role in a community, for example, an Aboriginal elder, a minister of religion;

(e) at least one lawyer, where possible one who is not engaged to advise the institution; and

(f) at least two people with current research experience that is relevant to research proposals to be considered at the meetings they attend. Such members may be selected, according to need, from an established pool of inducted members with relevant expertise.”

- Members will attend continuing education and training in research ethics at least every three years (National Statement 5.1.28 (b)).
• Membership appointments to the HREC will be considered for review every three years (National Statement 5.1.34). Recommended reappointments of individual members will be made by the HREC Chairperson to the Chair of Metro South Research for approval.

• Prospective members of the HREC may be recruited by direct approach, nomination or by advertisement.

• Members will be provided a letter of appointment including the date of appointment, length of tenure, assurance that indemnity will be provided in respect of the conduct of their duties as an HREC member, HREC meeting attendance responsibilities and general responsibilities of the position.

• As a contingency measure to address the COVID-19 Global Pandemic, MSHHS is collaborating with HIRO and other QH HRECs throughout Queensland to provide for business continuity and minimize disruption. HREC members will be asked to voluntarily review applications on behalf of other QH HRECs from across the state of Queensland. Members should be aware of this temporary additional scope and provide assistance and cooperation to ensure that the impact of COVID-19 on vital research is minimized.

4. Confidentiality and Conflict of Interest (National Statement 5.4.5)
Members will be required to sign, as part of their appointment process, a declaration form undertaking that:

• Any conflicts of interest, which exist or may arise during tenure on the Metro South HREC will be declared, and

• All matters of which they become aware during tenure on the Metro South HREC will be kept confidential.

5. Induction, Mentoring and Training
New members are given Induction training by staff from the Metro South Research office in conjunction with provision of written training material relevant to the position. They also attend initial meetings as an observer, and are provided individual mentoring, where possible, via the Chair and from time to time another member/s of the HREC.

All members attend continuing education courses and training in research ethics and regulation (at least every three years). Throughout their tenure, members are given the opportunity to attend conferences and workshops, supported by the Institution, that are relevant to the roles and responsibilities of the HREC.

6. Remuneration
Committee Members provide their services and expertise on a voluntary basis. All essential and necessary expenses incurred by members in carrying out their HREC duties will be reimbursed. Both the Chair and Deputy Chair/s of the HREC receive remuneration to compensate for the additional time required to chair the HREC meeting, direct low risk review outcomes and perform executive duties.

7. Committee Procedures
The Metro South HREC operates in accordance with written Metro South Research standard operating procedures (SOP). These procedures are reviewed at least every three years and updated as required. All HREC members have access to copies of the SOP, and from time to time, will be consulted regarding amendments.

(a) Protocol Submissions
• The HREC requires submissions to be in a standard format using the Human Research Ethics Application (HREA) Form (or its successor), which must be submitted via the Ethical Review Manager (ERM) website. The Committee requires researchers to electronically upload all supporting documents onto the ERM website.

• The Chair along with Committee members will determine if any additional expert advice is required in relation to scientific review.

Before giving a favourable opinion, the HREC should be adequately reassured that the research application meets the requirements of the Elements of Research set out in Chapter 3.1:

• Research Scope, Aims, Themes, Questions and Methods
• Recruitment of Participants
• Informed Consent
• Collection, Use and Management of Data and Information
• Communication of Research Findings or Results to Participants
• Dissemination of Research Outputs and outcomes
• After the Project

All submissions should adhere to the values and principles of ethical conduct as described in the National Statement on Ethical Conduct in Human Research (2007, updated 2018):

• Research merit and integrity
• Justice
• Beneficence
• Respect

(b) Levels of Ethical Review

• **Low & Negligible Risk Research (National Statement Section 2.1 & Section 5.1.18-5.1.23)**

Research that carries only negligible risk and involves the use of existing collections of data or records that contain only non-identifiable data about human beings may be exempted from full ethical review. The National Statement (2007, updated 2018) recognises that the levels of ethical review for low risk and negligible risk research may include, but need not be limited to:

a. review or assessment at departmental level by the head of department;

b. review or assessment by a departmental committee of peers (with or without external or independent members);

c. delegated review with reporting to an HREC; or

d. review by a subcommittee of an HREC.

In keeping with the National Statement, the Metro South HREC provides review of low risk research proposals via a Low Risk Review Panel comprised of the Low Risk Chair (fulfilled by the Committee Deputy Chair) and two nominated HREC members with expertise/understanding relevant to the nature and scope of the research and participants to be recruited. The Low Risk Chair will recommend approval of the low risk proposal, to be included on the agenda of the next HREC meeting. The HREC reserves the right to ratify the previous decision, instigate its own review process, request amendments, or clarification, or reject the proposal.
Research Identified as Greater than Low Risk  Any research identified as involving more than low risk must be referred to the HREC for full review, except in exceptional circumstances as stated below:

1. Multi-centred research studies reviewed and approved by another HREC  
The HREC, via the Chair, may approve a proposal with only expedited review, which another HREC has approved and the proposal conforms to the requirements of the Metro South HREC. The HREC reserves the right to ratify the previous decision, instigate its own review process, request amendments, or clarification, or reject the proposal.

The institution will accept the reviews of other QH Certified HRECs and those from Interstate health organisations identified as part of the National Mutual Acceptance.

2. Exceptional circumstances exempt from full ethical review  
In exceptional circumstances, where as a matter of public policy, and in the national interest, it is essential that an application should be reviewed urgently (to allow health-related research to commence as quickly as possible), the Metro South HREC Chair or Delegate may grant ethical clearance under exceptional circumstances for a protocol where:

- Another Committee has approved the protocol and the protocol conforms to the requirements of the HREC; and
- Clinical need necessitates urgent approval of the protocol.

(c) Meetings (National Statement sections 5.1.37 and 5.2.30-33)
- Meetings will be held monthly, except for January when there will be no scheduled meeting. However, in months involving significantly higher submission numbers a second meeting may be held following adequate period of notice to members (minimum two weeks).
- Meeting dates are available on the Metro South Research - Research Ethics and Governance website.
- Notice of meetings will be given to members for the current year and at least two (2) weeks prior to a meeting.
- A copy of the agenda, previous minutes, along with any other papers relevant to the meeting, will be forwarded to all members two weeks prior to the meeting.
- In addition, documents relevant to all new applications will be made available by the HREC Administration team, within the ERM Portal. The team will alert Committee members when documents have been made electronically available for access and review two weeks prior to the meeting. Members are then asked to submit, to the ERM Portal, their assessment/comments in respect to all new applications; prior to the close of business the day before the meeting. These comments are collated by the HREC Administration team and incorporated into a final set of HREC minutes.
- Meetings of the Low Risk Review Panel sub-committee will occur throughout each month of the year electronically via the ERM portal. Members will be rostered on for one month of the year and are expected to be available for the duration of the month to read and review submissions as allocated. Submissions are allocated via ERM and an email is sent out weekly to rostered members with the details of the studies and the review due date. Once allocated, members are expected to record their review comments in ERM within seven (7) days of allocation. If a member is unable to complete their review by the due date the
HREC Administration team must be notified as soon as possible so the review can be reallocated in a timely manner.

(d) **Meeting Protocol (National Statement sections 5.2.2-5.2.4, 5.2.28 – 5.2.33)**

- Decisions by the Committee as to whether the research protocol meets the requirements of the National Statement must be informed by the exchange of opinions from each of the members that constitute the minimum membership of the HREC.

- Where there is less than full attendance of the minimum membership at a meeting (quorum is one representative from each category as per 5.2.30), the Chair must be satisfied, before a decision is reached, that those members unable to attend the meeting have received all papers and have had an opportunity to contribute their views and that the views of all members have been recorded and considered. Members who are unable to attend a rostered meeting are asked to contribute and advise their opinion via ERM submission to the HREC Coordinator prior to the meeting.

- Meetings are held in the Translational Research Institute on Level 2, Princess Alexandra Hospital campus. Teleconference linkage to individual members unable to be present in person will be acceptable, if required.

- The Principal Investigator or a representative for the Investigator may be invited to attend the relevant meeting to discuss a proposal but would be required to leave the meeting before any decision is taken.

- Members of the Committee will be required to declare any conflict of interest (real or perceived) prior to or at any time during a meeting, such as when the member is associated with a research protocol under review by the Committee. The Committee will determine the action to be taken including excluding the member from the meeting during deliberation of the particular protocol.

- In general, HREC decisions will be reached by a consensus rather than simple voting majorities.

- The appointed Chair will chair every meeting when present. When the Chair is absent or excluded because of a conflict of interest, the Committee will appoint a Chair from attendees at the meeting if a Deputy Chair is unavailable.

(e) **Secretariat Support**

Secretariat support will be provided by staff of Metro South Research, herein named the HREC Administration team.

(f) **Decisions from HREC meetings**

Minutes will record major issues discussed, concerns expressed, decisions taken and reasons for rejection or requirement for change to the protocol and where necessary link those reasons to the National Statement. Notification of the Committee’s deliberations will be made directly to the Principal Investigator (the HREC does not engage in direct communication with sponsors unless in exceptional circumstances and with the agreement of the Principal Investigator).

(g) **Post HREC Approval Process**

- **Site Specific Assessment (SSA)**
  The SSA is an institutional research governance requirement. It involves assessing the suitability of a site at which the research is being conducted and identifying whether the ‘actual’ and or ‘in kind’ resources required for the conduct and completion of the project can be met by the Metro South Hospital and Health Service.
can only commence at a site upon authorisation by the Metro South Chief Executive or Delegate (including sign-off of SSA).

- **Public Health Act (PHA) application**
  If the research involves access to confidential information held by Queensland Health then a PHA application may be required under Section 281 of the Public Health Act 2005. This is submitted through the Office of Health Innovation, Investment and Research by the researcher, for sign off by the Director-General / Director-General delegate, after HREC approval is given.

- **QCAT (Queensland Civil and Administration Tribunal) application**
  If the research involves persons with impaired decision making capacity, an application to the Queensland Civil and Administration Tribunal (QCAT) for approval to conduct clinical research may need to be submitted should the research meet the definition set out in the *Guardianship and Administration Act 2000*. This is submitted by the researcher, after HREC approval is given.

- **Progress Reports**
  Progress reports on all approved research protocols are to be submitted to the HREC via ERM at least annually. The progress report is due by the 30th April each year for the lifetime of the project. The Committee may request more frequent progress reports, primarily based on the level of risk associated with the particular research protocol.

- **Adverse Event Reporting**
  The reporting of adverse events must adhere to the guidelines included in the NHMRC *Safety Monitoring and Reporting in Clinical Trials involving Therapeutic goods 2016 and its supplementary guidance documents*.

(h) **Insurance and Indemnity for HREC members**
As per the National Statement on Ethical Conduct in Human Research Section 5.1.9 the Metro South HHS provides Committee members with indemnity under Queensland Government Insurance Fund insurance policy.

(i) **Storage of Data**
Data may be collected, stored or disclosed in three mutually exclusive forms:

- **Individually identifiable data**
  The identity of a specific individual can reasonably be ascertained. Examples of identifiers include the individual’s name, image, date of birth or address, unique unit record number (URN);

- **Re-identifiable data**
  Identifiers have been removed and replaced by a code, but it remains possible to re-identify a specific individual by, for example, using the code or linking different data sets;

- **Non-identifiable data**
  Data which have never been labelled with individual identifiers or from which identifiers have been permanently removed, and by means of which no specific individual can be identified. A subset of non-identifiable data is that which can be linked with other data so it can be known that they are about the same data subject, although the person’s identity remains anonymous.

The NHMRC National Statement on Ethical Conduct in Human Research (2007, updated 2018) avoids the term ‘de-identified data’, as its meaning is unclear. While it is sometimes used to refer to a record that cannot be linked to an individual (‘non-
identifiable”), it is also used to refer to a record in which identifying information has been removed but the means still exist to re-identify the individual. When the term ‘de-identified data’ is used, researchers and those reviewing research need to establish precisely which of these possible meanings is intended.

(j) The Human Research Ethics Application (HREA)
In keeping with the National Statement on Ethical Conduct (2007, updated 2018) the Metro South HREC uses the HREA application form as its core document for all levels of review of research protocols, completed online by the researcher and submitted to the HREC Coordinator via ERM. The HREA and its supporting documentation, such as the participant information sheet, consent form(s) and the protocol serve to inform the Committee regarding the ethical aspects and scientific merit of the protocol.

8. Monitoring (National Statement Chapters 5.5 & 3.3)
Both the Institution and the HREC act in accordance with the National Statement in relation to monitoring approved research and require the Principal Researcher (including Co-ordinating Principal Investigator for multi-centre research) to:

- Keep adequate records (hard copy and/or electronic) and provide access to the Committee when requested.
- Provide annual progress reports at intervals specified by the Committee and at completion of any research.
- Notify and provide reports, in a timely fashion, to the HREC of significant events, complications and protocol violations that occur at any time during the conduct of research, detailing the course of action taken. Where relevant, Principal Investigators will notify the outcome of monitoring visits by trial sponsors. In relation to sponsored clinical trials and investigator-initiated trials involving drug or device interventions the notification of adverse events should be in keeping with the NHMRC Monitoring Framework.
- Provide prospective advice of any proposed changes to be made to the protocol and obtain approval of these changes prior to implementation.
- Notify the Committee if the research is to be discontinued before the expected date of completion (detailing a justification for the termination of the trial, such as closure of the trial by the pharmaceutical sponsor).
- Notify the HREC of any complaints received from participants, staff, researchers or the community.
- Provide documents of the outcomes of the research to the HREC.

The HREC may:
(a) Request an interview with the researchers if required.
(b) Request access to research data and records (including consent documentation as part of a random audit).
(c) Request the opinion of external experts if considered necessary.
(d) Liaise with the local site Research Governance Office.

9. Complaints (National Statement Chapter 5.6)
In keeping with the National Statement and the “Australian Code for the Responsible Conduct of Research 2018” the institution recognises the Manager, Research Integrity and Compliance,
Metro South Research, as the “designated person” for handling of research complaints in consultation with the Chair of the HREC, including research misconduct.

- In the first instance all complaints will go to the Chair of the HREC via the HREC Coordinator for consultation with the HREC.
- Any complaints received by the researcher must be forwarded to the Chair of the HREC via the HREC Coordinator.
- Complaints on the process, conduct or decisions of the Committee should be made in writing to the Chair of the HREC via the HREC Coordinator. The Chair of HREC will determine action to be taken. This may necessitate a special meeting of the Committee, which may be called without the usual 14 day requirement for notice.
- All complaints will be acknowledged within seven (7) days. The complainant will be advised of the decision of the HREC within 30 days. If the complainant does not accept the decision of the HREC, the complaint may be communicated by the Chair to the Metro South Chief Executive for further consideration.
- Any concerns, complaints or allegations about the conduct of a research protocol will be recorded in a register and sent to the local site Research Governance Officer. Processing of research complaints regarding the HREC review process will be as per the Queensland Government Department of Health HREC Administrators SOP and will also be recorded in a register.

10. Amendment to the Terms of Reference
Terms of Reference may be amended from time to time by following the procedure below:

- The proposal must be in writing and circulated to all HREC members for their consideration to allow for the views of members to be discussed at the next scheduled meeting of the HREC for ratification.
- Proposed amendments may be made by a member of the HREC or Secretariat staff; in conjunction with the HREC Chair.
- The amended terms of reference will be sent by the Chair to the Director Research Development and Ethics for forwarding to the Metro South Chief Executive.

11. Review
The Terms of Reference will be reviewed by the Committee every three years at the first meeting of the calendar year, unless requested earlier by a member of the HREC or Secretariat in conjunction with the HREC Chair. The date of next review is January 2022.

Endorsed:

A/Prof Scott Campbell
Chair
Metro South Hospital and Health Service Human Research Ethics Committee
Metro South Health

Dated: 01 / 05 / 2020
KEY DOCUMENTS AND WEBSITES


Australian Code for the Responsible Conduct of Research (2018)

Note for Guidance on Good Clinical Practice (CPMP/ ICH / 135 / 95):

World Medical Association Declaration of Helsinki - Ethical Principles for Medical Research Involving Human Subjects:
http://www.wma.net/en/30publications/10policies/b3/

Health Innovation, Investment and Research Office, Qld Health:

Metro South Research, Qld Health: https://metrosouth.health.qld.gov.au/research

Metro South Research Policies and Procedures: