



Final approval for ethics application

You are reminded that this letter constitutes **ethical** approval only. **Ethics approval is one aspect of the research governance process.**

You must not commence this research project at any SA Health sites listed in the application until a Site Specific Assessment (SSA), or Access Request for data or tissue form has been authorised by the Chief Executive or delegate of each site.

20 October 2016

A/Professor Peter Catcheside
Sleep Health Service
Respiratory and Sleep Services
Repatriation General Hospital
DAW PARK SA 5041

Dear A/Professor Catcheside

The Southern Adelaide Clinical Human Research Ethics Committee (SAC HREC EC00188) have reviewed and provided ethical approval for this application which appears to meet the requirements of the *National Statement on Ethical Conduct in Human Research*.

Application Number: OFR # 55.16 - HREC/16/SAC/207

Title: An investigation of wind farm and traffic noise effects on sleep

Chief investigator: A/Professor Peter Catcheside

Approval Period: 20 October 2016 to 20 October 2019

The below documents have been reviewed and approved:

- General Research Application (Parts 1 & 2) v1.3 dated 16 October 2016
- PICF Wind Farm Study (Part 1) v1.3 dated 03 October 2016
- PICF Wind Farm Study (Part 2) v1.3 dated 03 October 2016
- Community Liaison Group Document dated 26 April 2016
- Endorsement Letter SALHN, Respiratory and Sleep Services dated 01 June 2016
- Endorsement Letter School of Computer Science, Engineering and Mathematics Services dated 03 June 2016
- Endorsement Letter School of Psychology dated 23 May 2016
- Epworth Sleepiness Scale dated 06 June 2016
- EQ-5D Instrument Generic Health Status dated 06 June 2016
- Insomnia Severity Index dated 06 June 2016
- Pittsburgh Sleep Quality Index dated 06 June 2016
- Supplemental Material Questionnaire dated 06 June 2016

The below documents have been noted:

- Catcheside Grant Proposal dated 06 June 2016
- Assessor Snapshot Report dated 06 June 2016
- NHMRC Funding Document dated 06 June 2016

TERMS AND CONDITIONS OF ETHICAL APPROVAL

As part of the Institution's responsibilities in monitoring research and complying with audit requirements, it is essential that researchers adhere to the conditions below and with the *National Statement chapter 5.5*.

Final ethical approval is granted subject to the researcher agreeing to meet the following terms and conditions:

1. The approval only covers the science and ethics component of the application. A SSA will need to be submitted and authorised before this research project can commence at any of the approved sites identified in the application.
2. If University personnel are involved in this project, the Principal Investigator should notify the University before commencing their research to ensure compliance with University requirements including any insurance and indemnification requirements.
3. Compliance with the *National Statement on Ethical Conduct in Human Research (2007)* & the *Australian Code for the Responsible Conduct of Research (2007)*.
4. To immediately report to SAC HREC anything that may change the ethical or scientific integrity of the project.
5. Report Significant Adverse events (SAE's) as per SAE requirements available at our website.
6. Submit an annual report on each anniversary of the date of final approval and in the correct template from the SAC HREC website.
7. Confidentiality of research participants MUST be maintained at all times.
8. A copy of the signed consent form must be given to the participant unless the project is an audit.
9. Any reports or publications derived from the research should be submitted to the Committee at the completion of the project.
10. All requests for access to medical records at any SALHN site must be accompanied by this approval email.
11. To regularly review the SAC HREC website and comply with all submission requirements, as they change from time to time.
12. Once your research project has concluded, any new product/procedure/intervention cannot be conducted in the SALHN as standard practice without the approval of the SALHN New Medical Products and Standardisation Committee or the SALHN New Health Technology and Clinical Practice Innovation Committee (as applicable) Please refer to the relevant committee link on the SALHN intranet for further information.

Kind Regards



A/Professor Bernadette Richards
Chair, SAC HREC