Austin Hospital

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AUSTIN HEALTH HUMAN RESEARCH ETHICS COMMITTEE ETHICAL APPROVAL FOR AMENDMENT

Dr Matias B Yudi Austin Health

01 October 2019 (Updated 2 October 2019)

Dear Dr Matias B Yudi

HREC Reference Number: HREC/47081/Austin-2018

Austin Health Project Number: ND 47081/2018

Project Title: SMART-phone Based Cardiovascular Risk Reduction Program in Patients Undergoing Treatment for BREAST Cancer [SMART-BREAST]: A Randomized Controlled Trial

I am pleased to advise that the above project amendment has **received ethical approval** from the Austin Health Human Research Ethics Committee (HREC). This HREC is organised and operates in accordance with the National Health and Medical Research Council's (NHRMC) National Statement on Ethical Conduct in Research Involving Humans (2007, updated 2018), and all subsequent updates, and in accordance with the Note for Guidance on Good Clinical Practice (CPMP/ICH/135/95), the Health Privacy Principles described in the Health Records Act 2001 (Vic) and Section 95A of the Privacy Act 1988 (and subsequent Guidelines).

Original HREC Approval Date: 17/01/2019

Ethical approval for this amendment now applies at the following sites:

Sites	
Austin Health	

Approved Documents:

The following documents have been reviewed and approved:

Document	Version	Date
Protocol	4	13 September 2019
Austin Health Participant Information Sheet and Consent	4	13 September 2019
Form		

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Site Specific Assessment:

A copy of this letter must be forwarded to all Principal Investigators at every participating site and must be submitted to the relevant Research Governance Officer at each site.

Conditions of Ethics Approval:

- You are required to submit to the HREC:
 - An Annual Progress Report (that covers all sites listed on approval) for the duration
 of the project. This report is due on the anniversary of HREC approval. Continuation
 of ethics approval is contingent on submission of an annual report, due within one
 month of the approval anniversary. Failure to comply with this requirement may
 result in suspension of the project by the HREC.
 - A comprehensive Final Report upon completion of the project.
- Submit to the reviewing HREC for approval any proposed amendments to the project including any proposed changes to the Protocol, Participant Information and Consent Form/s and the Investigator Brochure.
- Notify the reviewing HREC of any adverse events that have a material impact on the conduct of the research in accordance with the NHMRC Position Statement: Safety monitoring and reporting in clinical trials involving therapeutic goods November 2016.
- Notify the reviewing HREC of your inability to continue as Coordinating Principal Investigator.
- Notify the reviewing HREC of the failure to commence the study within 12 months of the HREC approval date or if a decision is taken to end the study at any of the sites prior to the expected date of completion.
- Notify the reviewing HREC of any matters, which may affect the conduct of the project.
- If your project involves radiation, you are legally obliged to conduct your research in accordance with the Australian Radiation Protection and Nuclear Safety Agency Code of Practice 'Exposure of Humans to Ionizing Radiation for Research Purposes' Radiation Protection series Publication No.8 (May 2005)(ARPANSA Code).

The HREC may conduct an audit of the project at any time.

Yours sincerely,

Alyaa Mohamed HREC Non-Drug Trials Officer

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