

Research Governance Submission Checklist (Accepting Applications)

This checklist is intended to be used to support research teams in compiling and completing Site Specific Authorisation / Site Authorisation for accepting applications under National Mutual Acceptance (NMA)

It includes required evidence of ethics submission and approval (PART B), Site Specific Authorisation Requirements (PART C)).and additional requirements for Tele-trials (PART D).

This checklist is intended to be used as a guide only. Submissions requirements will vary with each application, it is important to determine the specific requirements for each submission. Submission requirements, including meeting dates and deadlines are outlined in the <u>Alfred Health Ethics and Governance website</u>, if you are unsure of the submission requirements for your application, please contact the ethics and research governance office to access their assistance and support.

Further information about each requirement is hyperlinked within the checklist. Please refer to the following websites for more information.

- ERM website
- <u>Victorian Government Clinical Trials and Research</u> website

Instructions

- This checklist must be completed and emailed with the application to the <u>Ethics & Research</u> <u>Governance Office</u>
 - Against each supporting document please indicate the following

 - \circ \Box No: Document is required to be submitted and is still outstanding
 - INA: Document is not required to be submitted with the application
- The SSA form requires supporting documents to be uploaded in ERM

Part A: Project details and investigator contact details

	Coordinator/Requester	Principal Researcher
Name		
Email		
Department		
Telephone		

	Project Details
HREC Reference Number	
Local Project Number/Protocol Number	
Full Project Title	
Target Clinical Area(s) e.g. ICU	
Campus involved in the project	□The Alfred □Caulfield
	□Melbourne Sexual Health Centre
	□Monash Alfred Psychiatry Research Centre
	□ Sandringham □Other (Please specify)

	Local Sponsor Details
Type of Sponsor	□ Industry □ Collaborative Group □ Investigator Initiated
Name of Contact	

Part B: Evidence of Ethics Submission and Approval

The following documents are required to be submitted to demonstrate evidence of ethics submission and approval.

All documents that are submitted should be the latest version

Su	pporting Documents	Yes	Pending	NA
Eth	nics Approval and related Correspondence			
•	Copy of initial and subsequent Amendment (if any) Ethics Approval Letters/Certificates Decision-making correspondence between the Reviewing HREC and Co- ordinating Principal Investigator			
Do	cuments approved by the reviewing Ethics Committee			
• • •	The latest version of the core ethics applications documents submitted to the Reviewing HREC including: Human Research Ethics Application (HREA) <u>Victorian Government Clinical Trials and Research website</u> Victorian Specific Module (VSM) <u>VSM in ERM guide</u> <u>Victorian Government Clinical trials and research website</u> Protocol or Project Description			
	<u>Alfred Health</u> Ethics and Research Governance Office <u>Victorian Government Clinical Trials and Research website</u>			
•	Master Participant Information and Consent Form(s): Master Opt-out information Alfred Health specific Master Information Sheets,			
•	Alfred Health specific Master <u>Participant Information and Consent Form(s)</u> Telephone Script (if participants will be contacted by telephone and/or verbal consent sought)			
•	e-Consent including screen shot of all information available on electronic devices including PICF			
•	search involving participants that are unable to consent <u>Medical Treatment Decision Maker Checklist</u> Standard Operating Procedure for participant enrolment or notification Master Information Sheets			
Dru	 and/or Device Information <u>Therapeutic Goods Administration</u> (TGA) <u>Office of the Gene Technology Regulator</u> (OGTR) 			
•	Investigator's Brochure or Product Information Consumer Medicine Information			
•	Instructions for Use (for devices) TGA ARTG Public Summary (if drug/device is TGA-approved) Office of the Gene Technology Register (OGTR) Licence or Confirmation of Exempt Dealing (if the study involves a Genetically Modified Organism))			
• Stu	Information about system (refer to electronic systems section) Idies involving lonising Radiation Procedures:			
	 <u>Alfred Health</u> – Ethics and Research Governance Office <u>Victorian Department of Health</u> <u>Victorian Government Clinical Trials and Research website</u> 			
•	Alfred Health Medical Physicist's Report Alfred Health Victorian Medical Physics Risk Assessment Form – fully signed			

Victorian Government Clinical Trials and Research website (VSM) Screen shot of all information available on electronic devices Product Information for Apps including device system requirements Privacy, Data Security and Terms of Use Recruitment and Promotional Material Advertisements Flyers, posters, radio/television scripts, newsletter articles Social media advertisements and platforms to be used Social media advertisements and platforms to sterening questionnaire, consent form (refer to electronic systems section) Letters, emails to potential participants, phone script Webpage materials Referral form Substantial Referral form Data Access Policy (for registries/databases/CQRs) Questionnaires, interview guides Recording including methodology/ device used Data Collection sheets/Case report forms Data Cullection sheets/Case report forms Data Cullection approval for projects involving use of established databases Retention Participant retention materials Collection of Biological Samples or Use of Samples from a Biobank Alfred Health – Ethics and Research Governance Office NHMRC – National Statement on Ethical Conduct in Human Research Biobank or Samples Access Policy Ethically Defensible Plan (If the research is likely to generate any incidental and/or secondary findings for the participant and/or their genetic relative(s)) Approval from custodian of biobank	Ele	ectronic Systems and Apps			
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PART C: Site Specific Authorisation Requirements

Site Specific Authorisation submission requirements, including meeting dates and deadlines are outlined in the <u>Alfred Health Ethics and Governance website</u> and the <u>Victorian Government Clinical Trials and Research website</u>

The following documents are required to be submitted to obtain Site Specific Authorisation

Sup	oporting Documents	Yes	No	NA
Site	Specific Authorisation Submission to Alfred Health:			•
•	Site Specific Authorisation (SSA) Form			
Leg	al and regulatory documents			
•	Jurisdictional Legislative Requirements			
•	Quick Reference Table – assists in determining the correct agreement and inde	mnity to use	ć	
Res	search Agreements			
•	Clinical Trial Preparation Agreement			1
•	Medicines Australia Clinical Trial Research Agreement:			
•	Medical Technology Association of Australia (MTAA) Clinical Investigation			
	Research Agreement (Device trials)			
•	Alfred Health Investigator-initiated, company supported			
•	Monash Partners Research Collaboration Agreement			
•	Alfred Health Material Transfer Agreements			
•	Alfred Health Data Transfer Agreement			
•	Clinical Trial Preparation Agreement ("Pre-Nup")			
•	Student Placement Agreement			
•	Amendments or Addenda to Agreements			
•	Third Party Service Agreements			
•	Alfred Health Research Contract Checklist			
•	Equipment Loan Agreement			
Ind	emnity Insurance & compensation			•
•	Insurance Certificate			
•	Medicines Australia (commercially sponsored and collaborative groups only)			
	 Medicines Australia Standard Indemnity to Alfred Health for 			
	Pharmaceutical Trials Oct 2012			
	 Medicines Australia HREC review only Indemnity to Reviewing 			
	HREC for Pharmaceutical Trials Oct 2012 (copy of the already			
	signed document)			
•	Medical Technology Association of Australia			
	 MTAA Standard Indemnity Form for a Clinical Investigation to Alfred Health Apr 2010 			
	MTAA HREC Review Only Indemnity to Reviewing HREC for a Clinical			
•	Investigation HREC Review Only Apr 2010 (copy of the already signed			
	document)			
Rec	julatory documents	L		
•	eCTN Draft or TGA Acknowledgment			
•	TGA ARTG Public Summary			
•	Evidence of Alfred Health legal review for non-standard agreements or wording in contracts			
Alfr	ed Health Governance Endorsements	·		
•	Genetically Modified Organisms Advisory Committee /Monash University			
	Institutional Biosafety Committee			
•	Data Custodian approval for projects involving use of established databases			
•	Clinical Innovations Committee			
•	Research Product Introduction			

•	Digital Health review for research				
Us	e of Alfred Health Resources				
•	Use of Alfred Health Resources Form				
٠	Resource Centre Declarations				
•	Baker Heart and Diabetes Institute Imaging				
٠	Biomedical Engineering				
٠	Biostatistical Assistance				
٠	Cardiology Services				
٠	Clinical Trials Pharmacy				
٠	Data and Analytical Services				
٠	General Service Request Form				
He	alth Information Services				
•	Access to electronic and paper medical records				
•	IT account creation (For CRA & monitor)				
٠	Intensive Care Services				
•	Lung Function Services				
٠	Nursing Services				
٠	Pathology & Anatomical Pathology				
٠	Performance Analysis and Clinical Costing				
٠	Public Affairs				
٠	Radiology, Nuclear Medicine and Radiation Oncology				
٠	Sleep Laboratory				
Не	ad of Supporting Department or Program Approvals				
•	Head of Department sign off				
•	Emergency Services Research & Education Committee				
•	ICU Research Committee				
•	Trauma Research Committee				
Qualifications and Training					
٠	Investigator CV				
•	Evidence of Investigators professional registration				
•	Evidence of Investigators Good Clinical Practice (GCP) Training				
٠	Evidence of GMO training				
Fees and charges					
٠	Ethics & Governance Fee Payment Form				
٠	Detailed site budget				

PART D: Additional requirements for Tele-trials

Submission requirements for tele-trials are outlined in the <u>Victorian Government Clinical Trials and Research</u> website

Ethics Application Documents					
•	Amendment – details of the site PI				
Le	gal and regulatory documents				
•	Standard CTRA tele-trial subcontract (between Primary site and Satellite site) for each Satellite site				
Со	nsent				
•	Master Tele-trial Participant Information and Consent Form (based on the primary site PICF)				
•	Satellite site specific Participant Information and Consent Form				
Le	gal and regulatory documents	-			
•	Standard CTRA tele-trial subcontract (between Primary site and Satellite site) for each Satellite site				
•	Agreement between the sponsor and the Primary site – additional wording reflecting use of the Tele-trial model				
Stu	idies involving Ionising Radiation Procedures:				
٠	Radiation Medical Physicist Risk Assessment				
Qu	alifications and Training				
•	Supervision Plan (between Primary site and Satellite site) for each Satellite				
	site				
٠	Satellite staff - CV's				
٠	Satellite staff – Evidence of GCP Training				
٠	Satellite staff – Evidence of professional registration				
Oth	Other				
•	Evidence the satellite site Research Governance Office has been notified of				
	the pending submission				
•	Evidence the Principal Investigator has endorsed the conduct of tele-trials at the satellite site				

PART E Additional Documents

As applicable to the research project

Additional Supporting Documents