

PHKL REC RESEARCH APPROVAL APPLICATION (Please complete all sections and attach all support		
Section A. General Information		
(i) Full Research Title:		
(ii) Protocol Number (if applicable):		□ N/A
(iii) NMRR ID (if applicable):		
(iv) Research Type		
☐ Clinical	☐ Basic/ Biomedical	
☐ Health Management	☐ Public Health/ Epidemiology	
☐ Health System	☐ Health Social Science/ Behavioural	
Health Policy Research	Systematic Review	
Action Research	Case Study/ Report/ Clinical Audit	
(v) Research Subtype		
☐ Interventional Study: BA/ BE	Interventional Study: Clinical Trial	
☐ Interventional Study: Community Trial	Interventional Study: Quasi Experimental	
Observational Study: Basic/ Biomedical	Observational Study: Clinical Economics	
Observational Study: Clinical Epidemiology	Observational Study: Patient Registry/ Da	tabase
Observational Study: Questionnaire		
(vi) Therapeutic Area		
Accident & Emergency	Neurosurgery	
Anaesthesiology	Obstetrics and Gynaecology	
☐ Cardiology	Oncology	
☐ Dermatology	Ophthalmology	
☐ Diabetes Mellitus ☐ Endocrine / Metabolic	☐ Orthopaedics☐ Primary Care	
	☐ Rheumatology	
☐ Gastroenterology	☐ Surgery	
☐ Haematology	☐ Transplantation	
Hypertension	☐ Traumatology	
☐ Infectious Disease	Urology	
☐ Neonatology	☐ Nephrology	
☐ Neurology	Others	
Section B. Investigator Details		
Name of Principal Investigator:		
Tel:		
H/P Number:		
Email:		
Designation:		
Department:		
Institution:		
Good Clinical Practice Certification	Yes	



ist of Co-Investigators:			
Name	Designation	Institution	Signature
ction C. Project Information			
Project Summary			
ot >500 words including introductio	n, methodology, objectives and e	xpected research outc	omes
i) Research Objectives			
rimary Objectives			
econdary Objectives			
i) Time Frame (The start date of a	the study should he after ethics a	oproval)	
xpected Start Date :	(Day/Month/Year)	oprovar)	
xpected End Date :	(Day/Month/Year)		
esearch Duration :	(Months)		
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<i>(</i> •)	
(iv)	Research Funding
□ N	No Funding
	ndustrial sponsor
	Name of funder:
l	Amount of funding:
	Research Grant
	Name of funder:
	Amount of funding:
	Amount of funding.
Sect	ion D. Ethical Consideration
(i)	How many research subjects are you planning to enrol into your study?
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(ii)	Will you seek written consent from the research subjects before they are recruited into your study?
` '	☐ Not applicable
	Yes (Please attach Patient Information Sheet and Informed Consent Form)
(iii)	What are the benefits of the study to the research subjects?
(iv)	What are the potential risks and burdens to the research subjects?
(v)	Will the research subject(s) receive financial reimbursement or other compensations for
(٧)	
	participating in your research?
	■ Not applicable
	Yes. Please state the amount
	□ No
(vi)	Does this study have insurance coverage to cover potential injury to the participants?
	☐ Not applicable
	☐ Yes
	□ No
(vii)	Will participants' details be anonymised?
` ′	Yes
	No. Please explain why and what you will do to ensure participants' confidentiality is protected.
(viii)	Where will the data be kept? How long will the data be retained?



(ix) Do any of the investigators in this st	tudy have conflict of interest to declare?			
☐ No☐ Ves. Please state the conflict of inte	araat .			
Tes. Please state the conflict of the	Yes. Please state the conflict of interest			
Section E. Declaration				
<u>Declaration</u>				
<u>Deciai autori</u>				
I declare that the information in this application form is accurate to the best of my knowledge and belief and I take full responsibility for it.				
I understand it is my responsibility to obtain approval where appropriate from PHKL Research and Ethics Committee before the project takes place. I agree to inform PHKL Research and Ethics Committee of any variations to the research during the application period or during the conduct of my research.				
Principal Investigator:				
Name:				
Date:				
Section F. For Office Use Only				
Date of Received				
Received By				
Signature				
PHKL REC Ref No.				
Remarks				
Section G. Review by PHKL REC Chairma	an/ Deputy Chaiman			
Additional actions or information	Yes			
required?	□ No			
If Yes, please specify				
11 100, p.10000 0p00,				
Reviewed by:				
Name:				
Date:				



Section H. Supporting Documents

No.	Items	Document Name	Version Number	Version Date	Please mark (x) if applicable
1.	Study Protocol/ Proposal Protocol shall include detailed description on introduction, objective(s), methodology, data analysis plan and a Gantt chart * Compulsory for all studies				☐ Mandatory
2.	Investigator's CV * Compulsory for all studies				☐ Mandatory
3.	Good Clinical Practice (GCP) Certificate * Compulsory for clinical trials				☐ Applicable ☐ Not applicable ☐ To be submitted once available
3.	Patient Information Sheet and Informed Consent Form (English) * Required for projects involved human subjects				Applicable Not applicable To be submitted once available
4.	Patient Information Sheet and Informed Consent Form (Bahasa Malaysia) * Required for projects involved human subjects				☐ Applicable ☐ Not applicable ☐ To be submitted once available
5.	Patient Information Sheet and Informed Consent Form (Other applicable language) * Required for projects involved human subjects				☐ Applicable ☐ Not applicable ☐ To be submitted once available
6.	Questionnaire/ Surveys/ Interview * Required when applicable only				☐ Applicable ☐ Not applicable ☐ To be submitted once available
7.	Case Report Form/ Data Collection Form * Required when applicable only				Applicable Not applicable To be submitted once available
8.	Patient's Diary * Required when applicable only				Applicable Not applicable To be submitted once available
9.	Investigator's Brochure * Required for clinical trial only				Applicable Not applicable To be submitted once available
10.	Trial Insurance Certificate * Required for clinical trial only				Applicable Not applicable To be submitted once available
11.	Clinical Trial Agreement * Required for clinical trial only. Draft agreement is acceptable.				☐ Applicable ☐ Not applicable ☐ To be submitted once available



12.	Decision Letter from				Applicable
	other Ethics Committees/				☐ Not applicable
	Regulatory Authorities				☐ To be submitted
	* Including negative decision,				once available
	or any modifications/				
	amendments during initial				
	submission submitted to other				
	Ethics Committee and/or				
	Regulatory Authority				
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13.	Subject Facing Materials				Applicable
13.	* Required when applicable				☐ Not applicable
	only				☐ To be submitted
	Only				once available
14.	Other Relevant				
14.					☐ Applicable☐ Not applicable
	Documents for this Study				
	(please list down)				☐ To be submitted once available
	-				once available
Please	e submit a signed elec	tronic copy of the f	orm and all	supporting	documents to
	kl.rec@pantai.com.my.	1,7		11 3	
For fu	rther enquiries, kindly call the F	PHKL Research and Ethics	Committee Sec	retariat at 03-	2280 1320/ 1321
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