



MDMRCI and ARMCI Cluster Research Ethics Review Committee



CRERC FORM 3.1 APPLICATION FOR INITIAL REVIEW

To be filled by investigator

CRERC Protocol Number:	Click or tap here to enter text.	Submission Date:	Click or tap here to enter text.
Sponsor Protocol Number:	Click or tap here to enter text.	Date Received by CRERC:	Click or tap to enter a date.
Protocol Title:	Click or tap here to enter text.		

Type of Research	☐ Clinical Research	☐ Clinical Trial	☐ Laboratory Research
	☐ Genetic Research	☐ Socio-Behavioral	☐ Public Health
	☐ Others: Click or tap here to enter text.		

Protocol Version No. & Date		ICF Version no. & Date	
Submitted to SJREB for review?	☐ Yes, date submitted _____; SJREB Code No. _____ ☐ No		

Study Duration and no. of subjects allocated to site	Click or tap here to enter text.
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Sponsor	Click or tap here to enter text.
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Principal Investigator	Click or tap here to enter text.
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Cellphone Number:

Click or tap here to enter text.

Fax Number:

Click or tap here to enter text.

Institution

Click or tap here to enter text.

Are you an employee of the sponsor?

☐ Yes

☐ No

Did you do consultancy or part time work for the sponsor?

☐ Yes

☐ No

In the past year, did you receive P250,000 or more from the sponsor?

☐ Yes

☐ No

Other ties with the sponsor:

Click or tap here to enter text.

Ethical Responsibility and COI Statement

I hereby pledge to address all forms of COI that I may have and perform my tasks objectively, protect the scientific integrity of the study, protect all human participants and comply with my ethical responsibilities as Investigator.

PI Signature

Click or tap here to enter text.

Request for Waiver of Informed Consent

Is there a request for waiver of informed consent?

☐ Yes

☐ No

If yes, what is the justification?

- ☐ This research could not practically be done without a waiver of consent.
- ☐ The research presents no more than minimal risk to the participants
- ☐ The waiver will not adversely affect the rights and welfare of the subjects
- ☐ When appropriate, the participants will be provided with pertinent information after the study

Received by CRERC
Admin Secretary:

Click or tap here to enter text.

Date
:

Click or tap to enter a date.



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