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IRB PROTOCOL EVALUATION FORM* (Form 3.2)						
IRB Protocol No.		Date (D	D/M/Y):			
Protocol Title:		Spr	onsor:			
Principal Investigator:		Contact no./ Em	nail			
Adviser:		Contact no./ Er	mail			
Study Coordinator:		Contact no./ Er	mail			
Type of Study:				Review Status:		
() Intervention () Epidemiology () Observational study () Document review () Individual based () Genetic () Social Survey () Others, specify - Descriptive				() Full Board () Expedited		
Description of the Study in brief: Mark whatever applies						
() Randomized Drug () Use of Genetic Materials () Double blind () Medical Device () Multicenter study						
() Single blind () Observational	() Vaccine () Oper () Questionnaire () Globa	al protocol () Inve		() Sponsor Initiated Initiated		

To be filled out by IRB					
ASSESSMENT POINTS		Indicate if the Study Protocol contains the specified assessment point		Page and paragraph where is found	REVIEWER COMMENTS
1.SCIENTIFIC DESIGN	YES	NO	N/A		
1.1. Objectives					
Review of viability of expected output					
1.2. Literature review					
Review of results of previous animal/human					
studies showing known risks and benefits of					
intervention, including known adverse drug					
effects, in case of drug trials					
1.3. Research design					
Review of appropriateness of design in view					
of objectives					
1.4. Sampling design					
Review of appropriateness of sampling					
methods and techniques					
1.5. Sample size					
Review of computation of sample size					
1.6. Statistical analysis plan (SAP)					
Review of appropriateness of statistical					
methods to be used and how participant data					
will be summarized					
1.7. Data analysis plan					
Review of appropriateness of statistical and					
non-statistical methods of data analysis					
1.8. Inclusion criteria					
Review of precision of criteria both for scientific merit and safety concerns; and of					
equitable selection					
1.9. Exclusion criteria					
Review of criteria precision both for scientific					
merit and safety concerns; and of justified					
exclusion					
1.10. Withdrawal criteria					
Review of criteria precision both for scientific					
merit and safety concerns					
2. CONDUCT OF STUDY					
2.1. Specimen handling					
Review of specimen storage, access, disposal,					
and terms of use					
2.2. PI qualifications					
Review of CV and relevant certifications to					
ascertain capability to manage study related					
risks					
2.3. Suitability of site					
Review of adequacy of qualified staff and					
infrastructures.					
2.4. Duration					
Review of length/extent of human participant					
involvement in the study					
3. ETHICAL CONSIDERATIONS					
3.1. Conflict of interest					
Review of management of conflict arising					
from financial, familial, or proprietary					
considerations of the PI, sponsor, or the study					
site					
3.2. Privacy and confidentiality					<u> </u>

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Review of measures or guarantees to protect			
privacy and confidentiality of participant			
information as indicated by data collection			
methods including data protection plans			
3.3. Informed consent process			
Review of application of the principle of			
respect for persons, who may solicit consent,			
how and when it will be done; who may give			
consent especially in case of special			
populations like minors and those who are			
not legally competent to give consent, or			
indigenous people which require additional			
clearances			
3.4. Vulnerability			
Review of involvement of vulnerable study			
populations and impact on informed consent			
(see 4.2). Vulnerable groups include children,			
the elderly, ethnic and racial minority groups,			
the homeless, prisoners, people with			
incurable disease, people who are politically			
powerless, or junior members of a			
hierarchical group			
3.5. Recruitment			
Review of manner of recruitment including			
appropriateness of identified recruiting			
parties			
3.6. Assent			
Review of feasibility of obtaining assent vis à			
vis incompetence to consent; Review of			
applicability of the assent age brackets in			
children:			
0-under 7: No assent			
7-under 12: Verbal Assent			
12-under15: Simplified Assent Form			
15-under18:Co-sign informed consent form			
with parents			
3.7. Risks			
Review of level of risk and measures to			
mitigate these risks (including physical			
,psychological, social, economic), including			
plans for adverse event management; Review			
of justification for allowable use of placebo as			
detailed in the Declaration of Helsinki (as			
applicable)			
3.8. Benefits			
Review of potential direct benefit to			
participants; the potential to yield			
generalizable knowledge about the			
participants' condition/problem; non-			
material compensation to participant (health			
education or other creative benefits), where			
no clear, direct benefit from the project will			
be received by the participant			
3.9. Incentives or compensation			
Review of amount and method of			
compensations, financial incentives, or			
reimbursement of study-related expenses			
3.10. Community considerations			
5.10. Community Considerations			

Review of impact of the research on the community where the research occurs and/or to whom findings can be linked; including issues like stigma or draining of local capacity; sensitivity to cultural traditions, and involvement of the						
community in decisions about the conduct of study						
3.11. Collaborative study terms of reference Review of terms of collaborative study especially in case of multi-country/multi-institutional studies, including intellectual property rights, publication rights, information and responsibility sharing, transparency, and capacity building						
		DECOMME	NDATION			
RECOMMENDATION DECISION: Approval Major Revisions Minor Revision Disapproval						
COMMENTS: ☐ Informed Consent ☐ Study Methodology ☐ Inclusion/exclusion criteria ☐ Study Design/ sampling design ☐ Sample size ☐ Data collection/ data gathering instruments ☐ Data Analysis						
Reviewer's Name: Date:						
Signature:						