 UNIVERSITY OF Baguio	Reference No.: QF-QAO-016	Revision No: 00	Effectivity Date: March 1, 2020
	POLICIES, PROCEDURES AND GUIDELINES		


Title	
REC PROTOCOL NO	

UBRDC-RCR-022 Form 1. RESEARCH PROPOSAL ETHICS REVIEW FORM FOR RESEARCHES INVOLVING HUMAN PARTICIPANTS

To the ethics reviewer: Evaluate the presence of the important ethical considerations in the application by checking the appropriate column.

COMPONENTS	YES	NO	REMARKS
1.The researcher/lead researcher has adequate trainings on the design and methodology to be used.			
2. The researcher/lead researcher possesses the necessary background knowledge to conduct the study.			
3.The research can be completed successfully in the indicated time.			
4.The research design is appropriate to the study.			
5.The proposed methods for testing the stated hypothesis are appropriate.			
6.The data analysis procedures are appropriate.			
7.The study is NOT a duplication of a previous study.			
8. The data collection procedures respect the privacy of the participants.			
9. The data collection procedures guarantee anonymity of the participants (if the informants/participants opt for non-disclosure of identity).			
10.The data collection procedures guarantee confidentiality of information (if informants/participants opt for non-disclosure of information).			
11.The selection of participants ensures objective and non-discriminatory procedures.			
12. A provision is made that participation in the research is voluntary and that participants can withdraw anytime.			
13. There is a provision made for the participants to be informed of the results of the study.			
14.There are sufficient measures to minimize the risks identified in the study.			
15. There is evidence that the participants will not suffer long term physical, emotional and psychological harm as a consequence of participating in the research.			
16. The participants will NOT be spending or using their resources for the research.			
17. There is a risk management strategy to address the safety of participants in studies involving invasive nondiagnostic procedures such as venipuncture and incision			
18.There is evidence of the contribution of the study to improvement of life, systems and processes and most especially to the groups involved.			



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In general, the study is

Low risk

Medium risk

High risk

COMMENTS/SUGGESTIONS:

Action Taken

The study is approved to commence

The study needs minor revisions prior to commencement

The study needs major revisions prior to commencement

Evaluated by:

 (Signature over printed name)
 Member, REC

 Date

