

Internal IRB Approval Form for Human Subject Research

I. Primary faculty/staff responsible for conducting data collection/study:

Name: _____

Email: _____ Telephone: _____

2. Purpose for collecting the data; How will the data be used? _____

3. Describe the following aspects of the data collection and study.

method (e.g. surveys, focus groups, interviews, etc.) _____

location (e.g. on-campus, off-campus, online, in/out-of class) _____

subject or audience (e.g. student, faculty, staff) _____

period (start date, end date, and time(s)) for collecting data _____

4. Will identifiable data be collected of participants? Yes No

5. Will any type of compensation, payments, or gifts be given to the participants? Yes No

6. Attach the instrument (e.g. survey or interview questions) that will be used to collect data.

7. How will the personal data (if applicable) be kept confidential and secure during and after the study? _____

8. Statement of Confidentiality: All data shall be treated confidentially and not shared without the participant's prior consent. Do you affirm that you will have each participant to complete a consent form, prior to collecting information, and that evidence thereof is provided to the IRB Committee prior to collecting data from subjects? Yes No

Print Name (Faculty/Staff Responsible)

Signature

Date

For IRB Committee Use Only

IRB Committee Member Reviewer: _____

Committee Member's Questions, Concerns, Comments: _____

Approval: Yes No

Print Name (IRB Chair)

Signature

Date