Human Informed Consent Form

Instructions to the Student Researcher(s): An informed consent/assent/permission form should be developed in consultation with the Adult Sponsor, Designated Supervisor or Qualified Scientist.

This form is used to provide information to the research participant (or parent/guardian) and to document written informed consent, minor assent, and/or parental permission.

- When written documentation is required, the researcher keeps the original, signed form.
- Students may use this sample form or may copy ALL elements of it into a new document.

if the form is serving to document parental permission	i, a copy of any survey of questionnaire must be attached.
Student Researcher(s):	
Title of Project:	
I am asking for your voluntary participation in my scientify you would like to participate, please sign in the appr	nce fair project. Please read the following information about the project. opriate area below.
Purpose of the project:	
If you participate, you will be asked to:	
Time required for participation:	
Potential Risks of Study:	
Benefits:	
How confidentiality will be maintained:	
If you have any questions about this study, feel free to	contact:
Adult Sponsor/QS/DS:	Phone/email:
	ou decide not to participate there will not be negative consequences. may stop participating at any time and you may decide not to answer any
By signing this form I am attesting that I have read and to participate or permission for my child to participate	I understand the information above and I freely give my consent/assent
Adult Informed Consent or Minor Assent	Date Reviewed & Signed:(mm/dd/yy)
Research Participant Printed Name:	Signature:
Parental/Guardian Permission (if applicable)	Date Reviewed & Signed:(mm/dd/yy)
Parent/Guardian Printed Name:	Signature:

Human Participants Form (4)

Required for all research involving human participants not at a Regulated Research Institution. If at a Regulated Research Institution, use institutional approval forms for documentation of prior review and approval.

(IRB approval required before recruitment or data collection.)

Student's Name(s)	itle of Project	
Must be completed by Student Researcher(s) in collaboration with a large submitted my Research Plan/Project Summary which a the Research Plan/Project Summary Instructions.	addresses ALL areas indicated in the Human Participants Section of in my project or other documents provided to human participants. ined. red by the IRB.	
BELOW - IRB USE ONLY		
Must be completed by Institutional Review Board (IRB) after review of the research plan. All questions must be answered for the approval to be valid. (If not approved, return paperwork to the student with instructions for modifications.) Approved with Full Committee Review (3 signatures required) and the following conditions: (All 6 must be answered) Risk Level (check one): Minimal Risk More than Minimal Risk Approved with Full Committee Review (3 signatures required) and the following conditions: (All 6 must be answered) Risk Level (check one): Minimal Risk More than Minimal Risk No than Minimal Risk No to the student (Form 2): No to the student (Form 3): No to applicable (No minors in this study) Mitten Minor Assent required for minor participants: No to applicable (No minors in this study) Mitten Parental Permission required for minor participants 18 years or older: No to applicable (No minors in this study) Mitten Informed Consent required for participants 18 years or older: No to applicable (No participants 18 yrs or older in this study) IRB SIGNATURES (All 3 signatures required) None of these individuals may be the adult sponsor, designated supervisor, qualified scientist or related to (e.g., mother, father of) the student (conflict of interest). Lattest that I have reviewed the student's project, that the checkboxes above have been completed to indicate the IRB determination and that I agree with the decisions above. Medical or Mental Health Professional (a psychologist, medical doctor, licensed social worker, licensed clinical professional counselor, physician's assistant, doctor of pharmacy, or registered nurse) with expertise related to this project.		
Printed Name	Degree/Professional License	
Signature	Date of Approval (Must be prior to experimentation.) (mm/dd/yy)	
Educator		
Printed Name	Degree	
Signature	Date of Approval (Must be prior to experimentation.) (mm/dd/yy)	
School Administrator		
Printed Name	Degree/Professional License	
Signature	Date of Approval (Must be prior to experimentation.) (mm/dd/yy)	