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 experimentation.)

Melissa Ramkisson	Adolescent Education and Awareness about Prescription Opioids
Melissa Ramkisson	
Student's Name(s) Robert Hildebrand	Title of Project 516-561-4493 Hildebrr@vschsd.org
Adult Sponsor Must be completed by Student Researcher(s) in collaboration with the Adult Sponsor/Designated Supervisor/Qualified Scientist: 1. I have submitted my Research Plan/Project Summary which addresses ALL areas indicated in the Human Participants Section of the Research Plan/Project Summary Instructions. 2. I have attached any surveys or questionnaires I will be using in my project or other documents provided to human participants. Any published instrument(s) used was /were legally obtained. 3. Any published an informed consent that I would use if required by the IRB. 4. Yes No Are you working with a Qualified Scientist? If yes, attach the Qualified Scientist Form 2.	
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.)	
	red) and the following conditions: (All 6 must be answered)
The state of the s	nimal Risk More than Minimal Risk
2. Qualified Scientist (QS) Required: Ye	
3. Designated Supervisor (DS) Required: Ye	
4. Written Minor Assent required for minor participants:	
Yes No Not applicable (No minors in this study)	
5. Written Parental Permission required for minor participants:	
Yes No Not applicable (No minors in this study)	
6. Written Informed Consent required for participants 18 years or older:	
Yes No Not applicable (No participants 18 yrs or older in this study)	
Approved with Expedited Review (1 signature required). Study involves either of the following: Human participants will only provide feedback on project design/student-designed invention or prototype. etc., no personal	
data will be collected and there are no health or safety hazards.	
AND STORES AND	
Student is the only subject of the research and no more than minimal risk is involved.	
IRB SIGNATURES (All 3 signatures required unless expedited review checked above) 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).	
I attest 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.	
Amy Indellicati	Psy. D. School Psychologist
Printer Name	Degree/Professional License
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Signature Date of Approval (Must be prior to experimentation.)	
Educator	
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ANGELO STANCO	MS securary education
Printed Name	Degree ()
HNUELOHILL	MS .
Signature	Date of Approval (Must be prior to experimentation.)
School Administrator	
Robert Milan,	School Administrator
Printed Name 0	Degree/Professional License
Cotest Yelun'	11/19/2019
Signature Date of Approval (Must be prior to experimentation.)	