

Phone: 936.294.3621 Fax: 936.294.3622

## **Continuing Review Form**

I. Protocol Informat	tion			
SHSU Protocol #: Date of Report:				
*A. Research Title:				
B. Personnel				
1. Principal Investiga	tor (PI)			
*Name:		*University Status	/Title:	
*Department:		*College:		
*Phone Number:		*Email A	ddress:	
Mailing Address:				
	required when PI is a s			
Department:		College:		
Phone Number:		Email Add	lress:	
Mailing Address:				
2 Co Investigators as	nd any other Key Pese	arch Personnel (Co-PIs		
Name				Hairmaita Ctatas/Title
Name	Department	College	Email Address	University Status/Title
II. Amendments				
	g the addition of resear	-		
	-	longer associated with	•	
Name	Department	College	Email Address	University Status/Title



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Yes (Please list th	ne names of new pe	rsonnel)		
Name	Department	College	Email Address	University Status/Title
B. Have you had any	change in funding	or sponsorship for th	nis research that has not be	en reported to the SHSU
PHSC? Yes	No			
If yes, please specify	v:			
C. Are you requesting	g any changes to th	e research protocol?	Yes No	
If yes, please specify	7:			
D. Are you requestin	g any changes to th	e consent documents	s? Yes No	
If yes, please specify	and attach the revi	sed consent form:		
E. Are you requesting	g any changes to th	e HIPAA authorizati	on? Yes No	
If yes, please specify	and attach the revi	sed HIPAA form:		
F. Are you requesting	g any other changes	to the research?	Yes No	
If yes, please specify	v:			
III. Review Process	Determination			
The research was	originally reviewed	and approved by the	PHSC under Full Board pro	ocedures.
The research was originally reviewed and approved by the PHSC under Expedited procedures				
The research is permanently closed to the enrollment of new subjects; all subjects have completed all				
research-related	interventions			
The research rem	nains active only for	long-term follow-u	p of subjects. (Long-term	follow-up refers to
collection of dat	a on survival or dis	ease status. It does n	ot include data collection a	activities such as
interviews, clini	c visits, lab tests, et	c.)		
Participants have never been enrolled on this study, and no additional risks have been identified since				
the last approval.				



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## IV. Findings from This Research

A. Describe any prelimir	nary results or findings from t	this research, if available. If the	e preliminary results are
suggestive of one interve	ention being better or worse t	han other(s), please discuss w	hen further findings will be
available.			
B. If there are preliminar	ry results of this study, indica	te if there is a change in any o	of the following:
(1) Risks associated	l with the research: Yes	No	
(2) Potential for beau	nefit to be gained from the re-	search: Yes No	
(3) Alternatives to	subject participation in the re-	search: Yes No	
(4) Participant willi	ingness to continue participati	ing in the research: Yes	No
	h	Van Na	
_		YesNo	
If yes, please indicate ho	ow and when the findings wer	e shared?	
V Doutisin out Engelle	ant and Damasmanhias		
v. Participant Enrollm	ent and Demographics	Total number of participants	Total number from non-SHSU
	Total number of participants enrolled since initial (if first	Total number of participants enrolled to date (since initial	Sites (if 6+68 is the grant
A. AGE RANGE	year) or last continuing review	PHSC approval- over all years)	holder or lead institution)
	(most recent year)		,
Age: Newborn to 2 Years			
Age: 3 to 6 Years			
Age: 7 to 11 Years			
Age: 12 to 15 Years			
Age: 16 to 17 Years			
Age: 18 to 64 Years			
Age: 65+ Years			
TOTAL			



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## B. By population description, indicate the number of participants enrolled:

POPULATION DESCRIPTION	Number enrolled at SHSU since the initial (if first year) or last continuing review (most recent year)		Total number (by category) of participants enrolled since initial PHSC approval (total number of years)			
Total						
Special populations						
Mentally Disabled or Mentally Ill						
Decisionally Impaired						
K-12 Students						
SHSU Students						
SHSU Psychology Subject Pool						
Fetuses (prior to delivery)						
Pregnant Women when <u>pregnancy</u> is the primary focus of the research						
Prisoners are primary focus of research						
Prisoners are <i>incidental</i> (not focus of research)						
SHSU Employees						
Pregnant Women Primary (focus of research)						
Pregnant Women Secondary (not focus of research)						
Other						
DEMOGRAPHIC	cate the number Black, not of Hispanic origin	er of participant	White	ed: not of nic origin	Other or Unknown	Total
Females						
Males						
Unknown						
Total						
D. Informed Consent Process  Are you planning to enroll additional  If yes, please specify:			)			



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	E. Informed Consent in Other Languages					
Have any partic	ipants whose prir	mary language is not English been	approached to partici	pate and/or be enrolled		
in the research?	in the research? Yes No					
If yes, for these	subjects, in whic	h language(s) was the informed co	onsent process conduc	ted?		
F. Complaints						
Have any comp	laints been receiv	red about the research? Yes	No			
If yes, for each	complaint describ	be the substance of the complaint,	when it occurred, the	complainant's		
relationship to t	he study, and how	w the situation was resolved.				
G. Participation	Declined					
•		or parents, guardians, or legally a	ithorized representativ	ves for the subject)		
•	• ,	earch after being approached?		res for the subject)		
_	_	ers of persons recruited who were	<del>_</del>			
ii yes, piease pi	ovide total numb	ers or persons recruited who were	not enroned.			
H. Withdrawal						
	imanta duammad a	ut of the mesopoush often initial annul	Imant & marticipation	2 Vec No		
		ut of the research after initial enrol		<u> </u>		
If yes, please pr	ovide total numb	er of participants who have droppe	ed out and the reasons	, if Known.		
I. Safety						
•	study-related ser	ious adverse events (SAE) have oc	ecurred for this protoc	ol since its date of		
•	•	ious adverse events (SAE) have oc	ecurred for this protoc	ol since its date of		
1. How many approval by the	he PHSC?	ious adverse events (SAE) have od  Description of Incident &	•			
1. How many	•		ecurred for this protoc Foreseeability	ol since its date of  Resolution		
1. How many approval by the	he PHSC?	Description of Incident &	•			
1. How many approval by the	he PHSC?	Description of Incident &	•			
1. How many approval by the	he PHSC?	Description of Incident &	•			
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2. Protocol V	iolations: Summa	arize any major protocol violations that h	ave occurred during the past reporting	
period. (These	e should already	have been reported to the IRB per the SH	ISU prompt reporting policy)	
Date of Incident	Subject ID #	Description of Event	Corrective	
INVESTIGAT	OR'S ASSURA	NCE		
I certify that the information provided in this continuing review form is complete and correct. I understand that				
as Principal Investigator, I am ultimately responsible for the protection of the rights and welfare of human				
subjects and the ethical performance of the research. I agree to comply with all applicable SHSU policies and				
		ral, state and local laws. I also agree to th		
procedures, une	approducto rede	and to the transfer of the tra	• 10110 Willing.	
(1) The res	search will only l	be performed by qualified personnel as sp	pecified in the approved research	
application and/or protocol,				
(2) No cha	anges will be mad	le to the research protocol (except when	necessary to eliminate apparent	
	•	ne subject), or the consent process withou	•	
(3) Legally effective voluntary informed consent/assent will be obtained from all human subjects, unless				
this requirement is waived by the SHSU PHSC,				
	•	•		
(4) Unanticipated problems involving risks to subjects or others (UPIRSO), serious adverse events, and				
other reportable events will be reported to the SHSU PHSC in a timely manner.				
*Logratify th	nat I have comple	ted the required educational program on	athical principles and regulatory	
	•		emear principles and regulatory	
requirements in Human Subject Protections.				