

Nova Southeastern University Office of Clinical Research 3300 S. University Drive Fort Lauderdale, FL, 33328-2004

# **Protocol Approval Record**

# For Non-Federally Funded Human Subject Research

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2.	Complete this form prior to Clinical Trial Agreement negotiations.

1. Complete Section I to determine whether this form is required.

- 3. The Principal Investigator (PI) is responsible for the completion of this form, including obtaining all personnel and Deans' signatures.
- 4. Once completed, this form must be electronically forwarded to the Office of Clinical Research ocr@nova.edu

# Sec

ction I – Research Questions						
		the type of study in detail below, e.g., including study population, study design, what you will be testing s are being measured.				
2. H	Has this study a	already been registered on Clinicaltrials.gov?				
	No	Yes, NCT#				
3. V	Will the NSU IR	B be the IRB of record?				
	No	Yes				
If no,	, please indica	te the IRB of record.				

## **SECTION II – Protocol**

PI:					
College / Center / Depart	ment				
Phone ext	Email	(	Cell phone		
Protocol Title:					
Proposed Start Date:	End Date:	D	ouration:Y	ear(s) /Month	n(s)
Instrument of Award:	Contract Sub	Contract			
External Sponsor:					
Address:					
Contact Person	Phone		Email		
Date of Executed CDA		-			
Section III- Staffing					
Instructions: The individu	ual members of the Pro	ject Staff, a	is well as the Dea	in of College or Dept.	Chair for the faculty or staf
member's primary appoir	ntment must sign this s	ection. If t	he PI is a Dean o	Dept. Chair, then his	/her supervisor must sign
this section.	<del>,</del>		<del>_</del>		
Name	Role	% Effort	College/Dept	Personnel Signate	ure Dean/Dept. Chair or Supervisor (if applicable as described above)

Name	Role	% Effort	College/Dept	Personnel Signature	Dean/Dept. Chair or Supervisor (if applicable as described above) Signature

<sup>\*</sup>The PI and all individuals on the study team will receive a separate Conflict of Interest email via Cayuse. Please monitor your Clutter and Junk email just in case.

<sup>\*</sup>College space and equipment is to be approved by the Dean, and NSU Health space and equipment should be approved by Leonard Pounds, Vice President NSU Health.

### **Section VI - Commitments**

Instructions: PI should initial each statement.
I agree to conduct the study in accordance with the current protocol and will only make changes in the protocol
after notifying the sponsor and IRB, except when necessary to protect the safety, rights, or welfare of subjects.
I agree to personally conduct or supervise the described investigation(s).
(If applicable) I agree to inform any patients, or any persons used as controls, that the investigational product are
being used for investigational purposes and I will ensure that the requirements relating to obtaining informed consent in 21
CFR Part 50 and Institutional Review Board (IRB) review and approval in 21 CFR Part 56 are met.
(If applicable) I agree to report to the sponsor adverse experiences that occur during the investigation(s) in
accordance with 21 CFR 312.64. I have read and understand the information in the investigator's brochure, including the
potential risks and side effects of the investigational product.
I agree to ensure that all associates, colleagues, and employees assisting in the conduct of the study are informed
about their obligations in meeting the above commitments.
I agree to maintain adequate and accurate records in accordance with 21 CFR312.62 and to make those records
available for inspection in accordance with 21 CFR 312.68
I will ensure that an IRB that complies with the requirements of 21 CFR Part 56 will be responsible for the initial and
continuing review and approval of the clinical investigation. I also agree to promptly report to the IRB all changes in the
research activity and all unanticipated problems involving risks to human subjects or others. Additionally, I will not make
any changes in the research without IRb approval, except where necessary to eliminate apparent immediate hazards to
human subjects.
I agree to comply with all other requirements regarding the obligations of clinical investigators and all other pertinent
requirements in 21 CFR Part 312.

### **SECTION VII - Principal Investigator**

I certify that:

- the above information and content of the proposal are true, accurate and complete
- that the budget reflects all appropriate expense items
- and that the project will be performed in compliance with university and sponsor policies, if funded.
- I certify that I will obtain all necessary approvals. For example, human subjects or biosafety, if applicable to my project, prior to initiating any research activities.

Signature, Principal Investigator	Date

#### **SECTION IX - Dean of PI's College**

I certify that:

- Personnel, space, and facilities are available to conduct/support the project as proposed and PI/research personnel are appropriately qualified to conduct the work.
- The project is appropriate to the goals and objectives of the College Unit
- The budget is approved.
- The space (noted above) is approved.
- The sponsor's restriction or disallowance of F&A recovery for this submission, if applicable, is approved.

Signature, Dean, Unit Director, or authorized delegate as applicable	Date