

Rev 05/10/18 ems

REACTIVATION REQUEST for NUNM's Institutional Review Board (IRB)

- Studies with a Final Review Approval Letter signed <u>less</u> than six (6) months ago, may use this Reactivation of a Closed Study Request Form. However, studies with a Final Review Approval Letter signed <u>more</u> than six (6) months ago, must file a new study application.
- Principal investigators may request reactivation of a study using the Reactivation Request Form only if the study was
 closed in IRB compliance and with a signed and dated Final Review Approval Letter. If a study is closed by the IRB for
 non-compliance for any reason, the principal investigator must contact the NUNM IRB Chair, Rich Barrett, ND at 503552-1758 to determine the appropriate course of action for reactivation to be considered.

Instructions: Please answer each question. If a question is not applicable, please put N/A in the box.

1.0 Investigator Information

1.1 Principle In	vestigator (PI):				
1.2 PI Telephone Number: 1.3 PI E-mail:					
1.4 PI Fax Number:					
1.5 Co-Investigator Name(s) and Contact Info:					
		2.0 Study Information			
2.1 IRB Numbe	r:				
2.2 Study Title:					
2.3 Original IRI	B Approval Date: _				
2.4 Location(s)	of Study Activity:				
		3.0 Protocol Status			
3.1 Date Final Review Form was signed by the IRB:					
3.2 Have there l	been any enrollmer	nts, interventions/interactions since the closure date?			
Yes	No				
If yes, please explain:					
	_				

Reactivation Request Form Study Title:

PI:

IRB #:

Approval Date:



Approval Date:

Rev 05/10/18 ems

3.3 Have any data be	en obtained since the closure date?
Yes	No
If yes, please explain:	
3.4 Are you requesting	ng that data collected since the closure date be used in data analysis?
Yes	No
If yes, please explain:	(Note: This request for reactivation must be approved by the IRB prior to any use of the data.)
3.5 Have there been a	any other research activities not addressed above since the closure date?
Yes	No
If yes, please explain:	
	4.0 Justification
4.1 Please thoroughly	justify the need for reactivation of your study:
	5.0 Data Handling
5.1 Will you be using	a study key?
☐ Yes	□No
If yes, please initial to	attest that only one person will have access to the study key:
5.2 Will data	a be transmitted outside of NUNM?
☐ Yes	□No
If yes, please describe	plans for how the data will be transmitted, received, stored and returned/destroyed:
Reactivation Request For Study Title:	rm
PI:	
IRB #:	





5.3 Please attest to th	e following statements by initialing each:				
5.3.1 Hard copi	es of study data will be stored in a secure environment (locked cabinets in a locked office).				
5.3.2 Electronic data will be stored on password-protected computers.					
5.3.3 Only data	necessary to meet the specific aims of the study will be collected and analyzed.				
5.3.4 No data will be collected or shared until all necessary approvals and forms are signed and in place.					
5.3.5 The study	will remain open during the publication period.				
	6.0 Adverse Event or Unexpected Problem				
6.1 Have there been a	ny complaints from subjects in the past approval period?				
☐ Yes	□No				
If yes, please describe.					
6.2 Have there been a	any adverse events or unexpected problems in the past approval period?				
□Yes	□No				
	n detail and indicate when the IRB was notified of the event or problem. Indicate dates that porting Form(s) was (were) submitted to the NUNM IRB.				
6.3 Does the study ha ☐ Yes	ve a Data Safety Monitoring Board (DSMB)? □ No				
	the date of the last DSMB review. (Please note that investigators are required to submit				
	WINM IRB at the time they are made available to the investigator.)				
	7.0 Protocol Modifications and Revisions				
7.1 Do you want to ac	ld any new co-investigators to the study?				
☐ Yes	□No				
If yes, indicate who yo RCR and HIPAA train	u are adding, and submit their names on an updated IRQ form and include the ing dates for each.				
Reactivation Request For Study Title: PI: IRB #: Approval Date:	rm				





IRB #:

Approval Date:

7.2 Were there any d	eviations from the final approved protocol in the past approval period?	
Yes	No	
If yes, please summari	ze these deviations:	
7.3 Do you wish to su	bmit a protocol modification at this time?	
Yes	No	
If yes, please describe document, using track	the purpose and rationale for the changes and attach proposed changes in all applicable changes:	? IRB
	8.0 Consent Form	
8.1 Please attach a cop	y of your current consent form for renewal if you are enrolling new subjects.	
8.2 Is this the original	consent form or a revised form?	
Original	Revised N/A	
If revised, please prov form(s) and HIPAA fo	ide date of NUNM IRB approval for the revision. Electronically send the current consent $rm(s)$.	t
	9.0 Publications, Presentations and Recent Findings	
9.1 Have there been a	my presentations or publications resulting from this study since the closure date?	
Yes	No	
If yes, please attach a	copy of the abstract, or the publication, with this application.	
	ny significant new findings either from this study, or a related study (through a literal hat would have an effect on participants' willingness to continue participating in this	
Yes	No	
If yes, please describe	and cite references:	
9.2.1 If applicable	e, have participants been informed of the new findings?	
Yes	No	
	10.0 Conflicts of Interest and Commercialization	
Reactivation Request Fo Study Title: PI:	m	



	mber of the research team have a potential conflic and/or study outcome?	ct of interest with this study that could affect				
Yes	No					
If yes, please descr	ibe the conflict of interest here, and disclose it in the	consent form:				
10.2 Does the PI, (Helfgott Research	Co-I and other researchers have a current conflict Institute?	t disclosure form on file at NUNM /				
Yes	No					
10.3 If there are conflicts of interests, please describe the ways in which you have and will minimize harm to research subjects and/or the objectivity of research: (If not applicable, please write 'N/A'.) 11.0 Required Signatures						
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	Principal Investigator	Date				
	Clinical Investigator (if different than PI)	Date				
	Name of person completing this form	Date				
IRB USE						
Institutional Review ☐ Approved ☐						
Chair or Committee	e Member Name:					
Signature: Date:						

Reactivation Request Form Study Title: PI:

IRB #:

Approval Date: