

NCI ADULT/PEDIATRIC CIRB APPLICATION FOR CONTINUING REVIEW

Attachment_B17_Cont_Rev

APPLICATION COMPLETION DATE: _____

OMB# 0925-0753, Expiration Date: 07/31/2021

The purpose of the information collection is to conduct reviews of clinical trial studies. NCI guidelines mandate the participation of institutions in the CIRB for Network group studies. You are being requested to complete this instrument so that we can conduct activities involved with the operations of the NCI CIRB Initiative. Although your participation in Network group research and completion of the forms is voluntary, if you wish to participate in the CIRB, you must complete all questions on the form. The information you provide will be combined for all participants and reported as summaries. It will be kept private to the extent provided by law.

NOTIFICATION TO RESPONDENT OF ESTIMATED BURDEN

Public reporting burden for this collection of information is estimated to average 60 minutes per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control number. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to: NIH, Project Clearance Branch, 6705 Rockledge Drive, MSC 7974, Bethesda, MD 20892-7974, ATTN: PRA (0925-0753). Do not return the completed form to this address.

This application, when completed, contains information required by CIRB members to conduct a meaningful review of the study so answer each question as completely as possible. If an answer to any question cannot be provided, please provide an explanation for the missing answer. If you have any questions regarding the completion of this application, please contact the CIRB Helpdesk at 888-657-3711 or ncicirbcontact@emmes.com.

GROUP STUDY ID NUMBER: <u>\$\$Study ID\$\$</u>
STUDY TITLE: \$\$Study Title\$\$
PROTOCOL VERSION DATE: This application should be based on the current CIRB-approved Protocol Version Date. Please provide the protocol and the informed consent document with this Protocol Version Date.

STUDY CHAIR	
Name	\$\$Study Chair name\$\$, \$\$Study Chair Degree\$\$
Title	
Institution/Address	\$\$Study Chair Address\$\$
Phone Number	\$\$InvestPrimaryPhone\$\$
E-mail	\$\$InvestEmail\$\$
FAX Number	\$\$InvestFAX\$\$
Administrative	
Assistant Name	
Administrative	
Assistant E-mail	

Administrative		
Assistant Phone		
Number		
STUDY CO-CHAIR (If applicable)	
Name		
Title		
Institution/Address		
Phone Number		
E-mail		
FAX Number		
Administrative		
Assistant Name		
Administrative		
Assistant E-mail		
Administrative		
Assistant Phone		
Number		
CONTACT PERSON	(Person to contact with questions about this application)	
Name		
Title		
Institution/Address	3	
Phone Number		
E-mail		
FAX Number		
4.0 CIDD Ct d C		
1.0 CIRB Study S	otatus	
CIRB stud	vith a check mark the current study status as defined by the CIRB. Please not y status definitions differ from CTEP study status definitions. The CIRB defined for your convenience.	
	Active: The study has received full approval from CTEP and the CIRB, has be civated by the Cooperative Group, and the study is open to accrual.	een
	Initial Activation Date:	
be	Approved but Not Yet Activated: The study has gone through CIRB reviewen fully approved by the CIRB however it has yet to be activated by the Coopoup.	
	Temporarily Closed to Accrual: The study is not completed but is temporarily participants. Participants currently enrolled in the study continue to addy intervention and/or are being followed.	•
	Temporary Closure to Accrual Date:	

1.1.4	Temporarily Closed to Accrual and Intervention Suspended: The study is not completed but is temporarily not accruing participants. Participants currently enrolled have had study intervention suspended.
	Temporary Closure/Intervention Suspension Date:
1.1.5	☐ Closed to Accrual, Participants still Receiving Intervention: The study has permanently closed to accrual however enrolled participants are still receiving study intervention.
	Closure to Accrual Date: Number of participants still on study intervention:
1.1.6	☐ Closed to Accrual, Participants have Completed Intervention: The study is permanently closed to accrual and all participants have completed study intervention. Participants are either in the follow-up phase or have finished participation in the study.
	Closure to Accrual Date: Number of participants still in follow-up:
1.1.7	☐ Withdrawn: The study is withdrawn by the Study Chair prior to CIRB final approval or withdrawn prior to activation by the coordinating Cooperative Group. Once withdrawn, all study activity will be considered completed with the CIRB. If the study is reactivated, it will have to be submitted to the CIRB and reviewed as a new study.
	Withdrawal Date:
1.1.8	Completed: The study is considered completed with the CIRB only when it has finished its planned course and all of the following are true.
	 a. The study has been closed to accrual. b. All participants have completed study intervention. c. All participants have completed all follow-up activities. Yes No
	d. Analysis of the data is complete. Yes No e. The study has met its primary objectives and a final study report/publication has been submitted. Yes No If Yes, provide a copy of the final report/publication.
	If all of the above five questions have been answered "Yes", the study will be permanently closed with the CIRB. Please go to Section 2.0 and complete the rest of the form as a final report to the CIRB.
1.1.9	Administratively Completed: The study is considered administratively completed with the CIRB when it has been stopped earlier than planned and all of the following are true.
	 a. The study has been closed to accrual. b. Participants are no longer receiving study intervention. Yes No

			ed. \tag Yes	□No
	d. No further a	ctivity or data analys	es are being performe	d
			∐ Yes	∐ No
	with the CIRB. I	-	tudy was stopped earlie	idy will be permanently clo er than planned then compi
0 E	nrollment Information			
2.1	Accrual target:			
•	2.1.2 Total number of 2.1.3 Total number of 2.1.4 Total number of 2.1.5 Total number of chosen to withday Describe specific Projected Enrollment In convenience, we have retained	participants who comparticipants still in follower participants whose sturbands formation at Study Institute of the study	dy intervention was terminations: titutions ng so that you can easily	n: minated early or who have
T.	ARGETED/PLANNED ENRO			
E	thnic category	Carrio		
		Sex/G	ender	
		Females	ender Males	Total
H	lispanic or Latino			Total
	lispanic or Latino lot Hispanic or Latino			Total
N	•			Total
N E	lot Hispanic or Latino thnic Category Total	Females t population in terms of	Males race:	Total
N E	thnic Category Total 2.2.2 Describe the target	Females It population in terms of LLMENT: Number of Su	Males race:	Total
N E	thnic Category Total 2.2.2 Describe the target	Females It population in terms of LLMENT: Number of Su	Males race: bjects	Total
N E	thnic Category Total 2.2.2 Describe the target ARGETED/PLANNED ENRO	Females It population in terms of LLMENT: Number of Su	Males race: bjects ender	
N E	thnic Category Total 2.2.2 Describe the targeton ARGETED/PLANNED ENRO acial Categories merican Indian /Alaska	Females It population in terms of LLMENT: Number of Su	Males race: bjects ender	
R A	thnic Category Total 2.2.2 Describe the targetory ARGETED/PLANNED ENRO acial Categories merican Indian /Alaska	Females It population in terms of LLMENT: Number of Su	Males race: bjects ender	

White		
Racial Categories: Total of all Subjects		

2.3 Current Enrollment Information at Study Institutions
For your convenience, we have retained the NIH formatting so that you can easily include the information in this application.

	Sex/Gender					
Ethnic Category	Females	Males	Unknown or Not Reported	Tota		
Hispanic or Latino						
Not Hispanic or Latino						
Unknown (Individuals not reporting ethnicity)						
Ethnic Category: Total of All Participants*						
Racial Categories						
American Indian/Alaska Native						
Asian						
Black or African American						
Native Hawaiian or Other Pacific Islander						
White						
More than one race						
Unknown or not reported						

PART B. HISPANIC ENROLLMENT REPORT: (Cumulative)	Number of	Hispanics o	r Latinos Enrolled	I to Date
Racial Categories	Females	Males	Unknown or Not Reported	Total
American Indian or Alaska Native				
Asian				
Black or African American				
Native Hawaiian or Other Pacific Islander				
White				
More Than One Race				
Unknown or not reported				

July 2018

Racial (Categories: Total o **	of Hispanics or					
*These	e totals must agree.	**These totals mu	st agree.				=
2.4		study recruitme he plan to addre		sing compa	red to the inter	nded schedule	? If concerns
2.5	progressing co	ment to the ethn ompared to the ir , what is the plar	ntended sch	edule as de			
3.0 0	ther Study Info	rmation					
3.1	Have any findi Monitoring Bo	ings from this stu pard?	ıdy been pr	esented or	published othe	er than to a Da	ta and Safety
	Yes	□No					
	If yes, explain	and attach the pi	resentation	s or publica	ations		
3.2	participants' r would include	Chair's knowledgo isks and benefits any new informa Formation on alte	on this stu ation about	dy become the drugs	available since or procedures u	the last CIRB used in this stu	review? Thi
	∐Yes	□No					
	If yes, explain	and attach releva	ant docume	nts			
3.3	or administrat document, or	en any changes in cive updates to the study participant approval if this is	ne protocol, t questionna	Cooperativaires since	ve Group model the last continu	l informed con	isent
	Yes	□No					
	updates since	ist all changes, re the last continuin tinuation. Includ	ng review a	pproval or	initial review a	pproval if this	is the first
3.4		igator's Brochur approval if this is		_		itinuing reviev	v approval o
	Yes	□No		Not applica	able		
	Please provide	e the version date	e of the mos	st current I	B:		
3.5	Have there be	en any updates o	r changes s	ince the las	st continuing re	view approva	l, or initial

July 2018 Page **6** of **8**

review approval if this is the first review for continuation, to the financial conflict of interest

	development or coordination of the study?
	□Yes □No
	If yes, explain
	3.5.1 Do any of the updates or changes result in new or revised significant financial conflicts of interest as defined in the Conflict of Interest Policy for Cooperative Group Phase 3 Clinical Trials?
	□Yes □No
	If yes, please provide a copy of the Cooperative Group's management plan to address the new or revised conflicts disclosed in question 3.5.
1.0 Ac	lverse Event and Unanticipated Problem Information
4.1	How is the study monitored for safety? Data and Safety Monitoring Board (DSMB) Safety monitoring committee Not applicable, explain
	4.1.1 Date of last DSMB or safety monitoring meeting:
	Attach the current DSMB report supplied to investigators.
	4.1.2 Date/approximate date of the next DSMB or safety monitoring meeting:
4.2	Has a toxicity summary report been prepared for the study?
	☐Yes ☐No ☐ Not applicable
	If yes, attach a copy of the current toxicity summary report supplied to investigators.
4.3	Since the last continuing review approval, or initial review approval if this is the first review for continuation, have there been any incidents, experiences, participant complaints, or outcomes that indicate participants or others may be at greater risk of harm (physical or otherwise) than previously anticipated?
	□Yes □No
	If Yes, explain
4.4	Have there been any unanticipated problems since the last continuing review approval or initial review approval if this is the first review for continuation?
	□Yes □No
	If ves. has the unanticipated problem been reported to the CIRB?

disclosures of the Study Chair or any persons listed on the protocol who are involved in the

July 2018

	□Yes □No
	If No, please provide a description of the unanticipated problem and any corrective action planimplemented
4.5	Since the last continuing review approval, or initial review approval if this is the first review for continuation, has anything occurred to cause the risk-benefit assessment to change?
	□Yes □No
	If Yes, explain
Summar	y of CIRB-Requested Supporting Documents Required, if applicable
	Protocol upon which this application is based Informed consent document with the same Protocol Version Date as the protocol Presentations and publications for this study (Question 3.1)
	Relevant information relating to participants' risks and benefits (Question 3.2) Investigator's Brochure (Question 3.4)
	Management plan to address new or revised conflicts (Question 3.5.1) Current DSMB/safety monitoring committee report (Question 4.1.1)
	Current toxicity summary (Question 4.2)

Thank you for completing the NCI Adult/Pediatric CIRB Application for Continuing Review. Please submit the completed application and the required supporting documents via email to either adultcirb@emmes.com or pediatriccirb@emmes.com.

July 2018 Page **8** of **8**