OMB #0925-xxxx

# OHSR RESPONSE TO REQUEST FOR REVIEW OF RESEARCH ACTIVITY INVOLVING HUMAN SUBJECTS

Expiration Date: xx/xx/xxxx

FAX:				E	Exempt: #:	5469
To:	Montello, Micha	ael				
	NCI					
	EPN - Executiv	e Plaza North, 702	24			
From	: Office of Huma	ın Subjects Resear	rch (OHSR)			
The The Boa The	e data being collecte ard (CIRB). The da erefore, data are no	collected by the CIF ed is for the sole p at available is not in of being collected o	RB, there is no intent to userpose of fulfilling the mintended in any use other on human subject participed Cooperative Group clin	ssion o the Centi than to conduct I ants as part of th	ral Institutiona RB review of ne CIRB Initiat	I Review studies. iive. The
Origi	nal Request Receiv	ved in OHSR on:	10/19/2010			
Resp	onsible NIH Resea	rch Investigator(s):	: Michael Montello, No	CI		
OHS	R review of your re	equest dated Thu,	Oct 14, 2010 has determ	ined that:		•
	determination of N Involving Coded P on Engagement of	lot Human Subject rivate Information of f Institutions in Hur	n of human subjects do n is Research is based on or Biological Specimens' man Subjects Research ( THAT MAY ALTER THIS	the interpretation (OHRP, Revised (October 16, 200	of 45 CFR 46 d October 16, 8). NOTIFY C	under "Research 2008) and Guidance
	The activity is desi	ignated <u>EXEMPT</u> ,	and has been entered in THAT MAY ALTER THE	the OHSR datab	oase. <u>PLEAS</u>	
			s IRB review. Please for ormation in order to dete			
	Confidentiality Agr	reement				
	Reliance					
	Amendment					
	Other					
Not	a' .		Office	Person SPC	Admin As	sist. CB
gh Sig	arlotte Holden, ob		Acting Director, OHSE	R	10/25/2 Date	010
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	nan Subjects Data: ogic Material:	No		□1 □2 [	□3 □4 [	□5 □6

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## REQUEST FOR REVIEW OF RESEARCH ACTIVITY INVOLVING HUMAN SUBJECTS

INSTRUCTIONS: Please type directly on this form. You can expand the document if you need more space. If your research involves a survey or questionnaire, please attach it to this completed form.

Completed forms (with all required signatures) may be sent to OHSR by FAX (301-402-3443), email to ohsr\_nih\_ddir@od.nih.gov, or by mail (2C146). If you have any questions, call OHSR at (301) 402-3444.

Date:	October 14, 2010			W.	
To: O	FFICE OF HUMAN	SUBJECTS R	ESEARCH, Build	ding 10, Room 20	C-146
	Mike Montello (Signature)	<u> </u>	7.	10/15/1.	<u> </u>
Through	: Steve Friedman_ (Signature of appropriation)	priate Official	for IC, e.g., Lab/I	Branch Chief)	_
Protocol Titl	e: NCI Central Instit	tution Review I	Board (CIRB)		
	I Principal Investig		4000 - 1000 1000 1000 1000 1000 1000 100	D, MBA	
9.			1,61		
IC: NCI/NIH	Laboratory/Bra	inch CIB/CTEF	/DCTD	25	
Building & R	loom No.: EPN, Rm	7036_ Tel. No:	301-435-9206 F	FAX No.: 301-48	0-4663
Is the Princi If no, please	pal Investigator an l explain:	NIH employee	? _XYes _	No	Э

1. What is the proposed research activity that you intend to perform at NIH (please use lay terms):

Though data is being collected by the CIRB, there is no intent to use this information for research in any form. The data being collected is for the sole purpose of fulfilling the mission of the Central Institutional Review Board (CIRB). The data available is not intended for any use other than to conduct IRB review of studies. Therefore, data are not being collected on human subject participants as part of the CIRB Initiative.

The CIRB improves access to NCI-sponsored Cooperative Group clinical trials for potential study participants and their physicians by enabling local IRBs to rapidly approve clinical trials through the use of a facilitated review process. The NCI Central Institutional Review Board (CIRB) Initiative reviews protocol, informed consent document(s), completed CIRB application(s); and when appropriate, an investigator drug brochure. The CIRB reduces the administrative burden on local IRB's and investigators.

2. If applicable Name	le, list your non-NIH Collab Institution	Address Tel. # FAX #
	RB began meeting January 20	Research is not being conducted. 01; The Pediatric CIRB began meeting
Proposed com	pletion date? There is no re	esearch being conducted.
4. Will you be	these samples or dat	a?
Collecting Receiving Sending	Yes/No	
(b) Or are Yes	y exist?Yes _X_No	express purpose of this study?
(c) Or a co	ombination of (a) and (b)?	YesNo earch project? (Check all that apply)
Consultan Author of (identified in co- Co-author You or NI Decisional approved site?	question #2). ship on publication(s)/manus IH hold an IND for this resear authority over the design or If so, please explain.	olemented by your collaborating investigator cript(s) pertaining to this research.
sponsored Coo	operative Group clinical trials enabling local IRBs to rapidl	et. The CIRB improves access to NCI- s for potential study participants and their y approve clinical trials through the use of a

### 7. Where are the subjects of this research activity located?

Though data is being collected by the CIRB, there is no intent to use this information for research in any form. The data being collected is for the sole purpose of fulfilling the mission of the CIRB. The data available is not intended for any use other than to conduct IRB review of studies. Therefore, data are not being collected on human subject participants as part of the CIRB Initiative.

3. If human subjects are located elsewhere (not at NIH), w	vill you have direct
contact or intervention with them? (Examples: as subject's	physician; in obtaining
samples directly from the subject; by interviewing the subject	?)Yes No

#### \* no human subjects

9. What kind of human samples (e.g., tissue, blood) or data (e.g., private information, responses to questionnaires) will be involved in your research?

The NCI uses various information collection tools to support CIRB activities, though none of them are being collected for research purposes. These include forms requiring completion by a site conducting a clinical trial eligible for review by the CIRB, as well as forms completed by the CIRB members themselves.

There are 6 general areas in which forms are completed and data are collected. These include:

- CIRB Helpdesk survey: collects customer feedback pertaining to the use of the CIRB helpdesk.
- CIRB Institution Enrollment Forms: Enrollment documents collect information on the institutions interested in enrolling or having already enrolled in the CIRB Initiative.
- CIRB Membership Information: Membership documents collect information on CIRB members. Information requested includes: name, degree, college and/or university attended, graduation year, bio sketch, headshot, title, place of employment, address, email address, telephone number, and picture. This is public accessible information.
- Direct Deposit Form: This is an optional form for CIRB Members to complete should they choose to receive honoraria via direct deposit.
- IRB Application Forms: The application forms are required to be submitted to the CIRB in order to complete a submission for review.
- 6. <u>CIRB Reviewer Forms:</u> Theses forms are completed by the Board Members when they submit reviews.

3

The CIRB Helpdesk Survey seeks to collect information that can be used by the CIRB in a systematic fashion to improve operations and enhance reviewer satisfaction with the initiative. The remaining forms are necessary in order to facilitate a complete review of protocols by the CIRB with the absence of conflict of interest in place.

Last revised 8/4/09

10. If the samples, data do not come from an IRB approved protocol, do they come from:
(a) Repository Yes X No
(b) Pathological waste Yesx No
(c) Autopsy material Yesx No
(d) Publicly available sourceYes X_ No
(e) Other
Cooperative Groups, Board Members, and Institutions supply a variety of information for the conduct of the CIRB, though none of the data collected is for the intent of research.
11. Please check the box(es) that apply(ies) to the samples/data that you will receive.
(a) Samples and/or data will be anonym zed/unlinked. (The samples/data cannot be linked to individual subjects by you or your collaborators at other sites.)
(b) Samples and/or data will be coded, however that code cannot be used by either the sender or the receiver to identify specific individuals.
(c) Samples and/or data will be coded so that the provider of the samples/data can link them to specific individuals but the receiver will not be able to do so.
12. Will you send results back to the provider(s) (listed in question 2 of this form)?
(a) No, I will not send results back to the provider(s).
(b) Yes, I will send aggregate results to the provider(s).
(c) Yes, I will send results to the provider(s) that are linked to identifiable individuals.
If yes, does the provider intend to link your data to identifiable individuals?  Yes  No
13. Has the research activity that you are proposing in this form been approved by an Institutional Review Board (IRB) elsewhere?
There is no research being conducted. This project is Exempt. The reason for the exemption would be as cited in 45 CFR 46.101(b) (4): (4) Research involving the collection or study of

Last revised 8/4/09

existing data, documents, records, pathological specimens, or diagnostic specimens, if these

sources are publicly available or if the information is recorded by the investigator in such a manner that subjects cannot be identified, directly or through identifiers linked to the subjects.

	een reviewed by the following IRB (s)
Address Name of Title of	of institution that provided the review as of reviewing institution of PI for the IRB approved protocol f IRB approved protocol # al Wide Assurance (FWA) number**
No IRB review of the research activity desc	ribed in question #1 above has taken
(**An FWA is a contract between the U.S. Department of DHHS) and an entity receiving DHHS funds to will follow ethical guidelines and federal regulate subjects. For a list of domestic and international antip://ohrp.cit.nih.gov/search/asearch.asp#ASUR	conduct clinical research that the latter ions for the protection of human institutions go to
14. Per NIH guidance***, have conflicts of interesolved? _x_YesNo	rest by NIH employees, if any, been
f your answer is no, please see your Clinical E proceeding with this research.	Director about this matter before
***The January 5, 2005 NIH Guide to Preventing research conducted at NIH, <a href="http://ohsr.od.nih.gov">http://ohsr.od.nih.gov</a>	

## Brentin, Christine (NIH/OD) [E]

From:

Jennifer Dugan [jdugan@emmes.com]

Sent:

Tuesday, October 19, 2010 10:07 AM

To:

OHSR (NIH/DDIR); Pursley-Crotteau, Suzanne (NIH/OD) [E]

Cc:

Montello, Mike (NIH/NCI) [E]; Friedman, Steve (NIH/NCI) [E]; Adler, Jeanne (NIH/NCI) [E]; Valmonte,

Claudine; Goldberg, Jacquelyn (NIH/NCI) [E]

Subject:

OHSR Application

Follow Up Flag: Follow up Flag Status:

Red

Attachments:

OHSR App\_101510.pdf

Dear OHSR,

Please review the attached CIRB Application.

Per instruction, signatures were obtain in advance and hopefully any remaining items can be resolved via email.

Thanks,

Jennifer

Jennifer L. Dugan, MS Project Manager NCI CIRB Operations Office The EMMES Corporation 401 N. Washington Street, Suite 700 Rockville, MD 20850 Phone: 301-251-1161 x2827 Fax: 301-251-1355 jdugan@emmes.com www.emmes.com

## OHSR (NIH/DDIR)

From:

OHSR (NIH/DDIR)

Sent:

Tuesday, October 19, 2010 2:35 PM

To:

Montello, Mike (NIH/NCI) [E]

Cc:

'jdugan@emmes.com'

Subject: Request for Review Rec'd-OHSRP 5469

Good afternoon Dr. Montello,

This email is to verify that OHSR has received your Request for Review of Research and it is currently being processed as OHSRP #5469. Please use this number in any future correspondence regarding this study. We will contact you via email if any additional information is needed. If you have not heard from OHSR within 7 business days, please contact us.

Protocol Title: NCI Central Institution Review Board (CIRB)

Thank you.

Sincerely,

OHSRP - National Institutes of Health

Bldg 10, Suite 2C146 Bethesda, MD 20892

Office Telephone: 301-402-3444

Office Fax: 301-402-3443

The NIH is committed to maintaining the highest standards for the protection of human subjects.



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