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

## STATEMENT OF CONFIDENTIALITY

Add Note

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 20 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. Add Note Please refer to the Quickquide on Completing the Study Specific Worksheet for further quidance. Add Note View Audit Reason for submission: (Required) Open New Study: This study is not opened at the Signatory Institution. This is the first submission to the CIRB of a Study-Specific Worksheet About Local Context for this study at this Signatory Institution. Revision: This study is already opened at the Signatory Institution. This is a revision to the existing Study-Specific Worksheet About Local Context for this study at this Signatory Institution. Next Save for Later More • Add Note View Audit **Signatory Institution Information Submitting User Information** Campbell, Brian Email: bcampbell@emmes.com Add Note View Audit Enter the Study ID Number. (Click here if you would like to review a list of studies currently covered by NCI CIRB) (Required) Add Note View Audit Signatory Institution (Required) CIRB Operations Office ▼ Calculated Field Add Note View Audit No answer provided. No answer provided.

Previous Next Save for Later More

General Information	Add Note	View Audit
	questing to onen this study	
<ol> <li>Enter the email address of the Principal Investigator who is re (Required)</li> </ol>	questing to open this study.	
	If the PI's name does not appear above the email address field, this mea no active account associated with this email address. Please confirm the address is correct and that it is the email address associated with the PI If the email address is correct and the PI name still does not appear, you contact your Signatory Institution's RUMS Update Person and request the added to the CIRB Roster in RUMS.	email in IAM. will need to
Previous Next Save for Later More		
Site Reviewer	Add Note	View Audit
No answer provided.		
The topics listed below reflect those asked on the Annual Pri been completed. Indicate for each topic whether or not there are changes, please describe. If any of the 'Changed' answer Question 33.	e are any changes from the information previously provided	If there
2. General Information (Questions 1-2 on the Annual Principal In	Add Note  Ivestigator Worksheet About Local Context)	View Audit
No Change     Changed		
3. Research Staff (Questions 3-5 on the Annual Principal Investig (Required)	Add Note gator Worksheet About Local Context)	View Audi
No Change Changed		
4. Principal Investigator Resources (Questions 6-7 on the Annual (Required)	Add Note I Principal Investigator Worksheet About Local Context)	View Audit
No change Changed		
5. Recruitment (Questions 8-9 on the Annual Principal Investigat (Required)	Add Note tor Worksheet About Local Context)	View Audi
○ No Change ○ Changed		
6. Compensation to Study Participants (Question 10 on the Annua (Required)	Add Note al Principal Investigator Worksheet About Local Context)	View Audi
○ No Change ○ Changed		
7. Informed Consent Process (Questions 11-20 on the Annual Pri	Add Note ncipal Investigator Worksheet About Local Context)	View Audi
○ No Change ○ Changed		
8. Pharmacy Information (Questions 21-22 on the Annual Princip	Add Note al Investigator Worksheet About Local Context)	View Audit
No Change Changed		

9. Measures to Protect Confidentiality (Question 23 on the Annual Principal Investigator Worksheet About Local Con (Required)	Add Note text)	View Audit
○ No Change ○ Changed		
10. Measures to Protect Privacy (Question 24 on the Annual Principal Investigator Worksheet About Local Context)  (Required)	Add Note	View Audit
O No Change Changed		
11. Emergency Resources (Question 25 on the Annual Principal Investigator Worksheet About Local Context)  (Required)	Add Note	View Audit
○ No Change ○ Changed		
12. Using a Legally Authorized Representative (LAR) (Questions 26-28 on the Annual Principal Investigator Worksho Context)  (Required)	Add Note eet <b>About Loc</b>	View Audit
○ No Change ○ Changed		
13. Vulnerable Populations (Question 29 on the Annual Principal Investigator Worksheet About Local Context)  (Required)	Add Note	View Audit
○ No Change ○ Changed		
14. Additional Confirmations When Investigator Intends to Enroll Pregnant Women [45 CFR 46.204 (h), (i), (j)] (Que Annual Principal Investigator Worksheet About Local Context)  (Required)	Add Note estions 30-32	View Audit 2 on the
○ No Change ○ Changed		
15. Additional Information (Question 33 on the Annual Principal Investigator Worksheet About Local Context)  NOTE: If there are any changes to the documents approved by the CIRB, use track changes to clearly identify the re track additional changes and not changes that are already part of your institution's approved boilerplate language.	Add Note	View Audit
(Required)  No Change Changed or New Information		
If 'Changed or New Information', describe changes or new information.	Add Note	View Audit
ASC		

If any of the 'Changed' answers can be supported by an attachment, an attachment can be added here.	Add Note	View Aud
Add Attachment		
Additional Study-Specific Materials for Review (If Applicable)  Complete this section if you have any of the following study-specific materials to be reviewed by the CIRB.	Add Note	View Aud
16. Recruitment material(s).		
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	Add Note	View Audi
17. Assent form or consent at the age of majority form.	7100 71010	7,017,00
AQC.		
If applicable, an attachment can be added here.	Add Note	View Audi
Add Attachment		
<ul> <li>18. Translated documents for this study. Translated documents include, the institution's boilerplate language, shor form, or template document for consent at age of majority.</li> <li>Note: The following documents are required when submitting translated material:</li> <li>1. CIRB-approved English language document(s) corresponding to the translated document with a version or version.</li> <li>2. Translated version(s) of the CIRB-approved English language document with a version or version date that main translator's Certificate(s) of Accuracy or equivalent document(s) with reference to the version or version date.</li> </ul>	sion date tches the Eng	late assen
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If applicable, an attachment can be added here.	Add Note	View Audi
Add Attachment		
Study ID Number for the Transfer of Review Responsibilities from the CIRB to the new IRB for Study #2	Add Note	View Audi
Previous Next Save for Later More		
Form Completed You've completed the form. You can now either save the form for later revision, or submit it.		
Go Back Save for Later Print Submit		