

OMB Text

National Cancer Institute
Central IRB Tritiative
5 - Unanticipated Problem and/or Noncompliance Form -- CIRB Form

Next

Add Note

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OMB#: 0925 - xxxx Expiry Date: xx/xx/xxxx

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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 (0925-0625). Do not return the completed form to this address.

Signatory Institution Information	Add Note	View Audit
Submitting User Information		
Campbell, Brian		
Email: bcampbell@emmes.com		
Name of Signatory Institution		Add Note
(Required)		
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General Information		Add Note
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General Information 1. Enter Study ID Number. (Click here if you would like to review a list of studies currently covered by NCI CIRB) (Required)		Add Note
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Section C: Potential Unanticipated Problem	Add Note
1. Is this incident, experience, or outcome unexpected?	
(Required)	
© Yes ◎ No	
If Yes, describe how the incident, experience, or outcome is unexpected and/or add an attachment.	Add Note
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Attachment:	Add Note
Add Attachment	
2. Is this incident, experience, or outcome related or possibly related to participation in the research? (Required)	Add Note
○ Yes ○ No	
If Yes, describe how the incident, experience, or outcome is related or possibly related to participation in the research and/or add an attachment.	Add Note
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Attachment:	Add Note
Add Attachment	
3. Did the incident, experience, or outcome place the study participant(s) or others at a greater risk of harm? (Required)	Add Note
© Yes ○ No	
If Yes, describe how the incident, experience, or outcome placed the study participant or others at a greater risk of harm and/or	Add Note
add an attachment.	Add Note
Attachment:	Add Note
Add Attachment	
4. Describe any action the Principal Investigator and/or Signatory Institution has taken, is taking, or is planning to take, to address the incident, experience, or outcome.	Add Note
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Add an attachment, if applicable.	Add Note
Add Attachment	