NCI CTRP Attachment 3c

NCI CTRP Amendment Portal Workflow and Screen Shots

Step 1: User accesses the NCI Clinical Trials Reporting Program website at http://trials.nci.nih.gov – see screenshot, page 2

Step 2: User enters "Username" and "Password" – see screenshot, page 2

Step 3: User reviews NCI Clinical Trials Reporting Program burden statement – see screenshot, page 3

Step 4: System displays "Search Submitted Clinical Trials" page – see screenshot, page 4

Step 5: User selects to "Submit Trial Amendment" and amends an existing trial record – see screenshots, pages 5 – 10

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CTRP Home and Login page Clinical Trials Reporting Program Login

Clinical Trials Reporting Program	You are accessing a U.S. Governme this computer, (2) this computer ne network, and (4) all devices and sto a computer on this network. This Government-authorized use only, system may result in disciplinary penalties. By using this information the following: You have no reasonable exp communications or data transiting any time, and for any lawful Gov monitor, intercept, record, and sea transiting or stored on this informa Any communication or data transiti may be disclosed or used for any la
 Cancer Institute onal Institutes of Health	POLICIES Accessibility Disclaimer FOIA Privacy & Security

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HHS Vulnerability Disclosure

HELP AND SUPPORT ctrp_support@mail.nih.gov

LAST BUILD 04-19-22 10:51

API Build Details

U.S. Department of Health and Human Services | National Institutes of Health | National Cancer Institute | USA.gov

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CTRP Burden Statement

Clinical Trials Reporting Program Registration

NCI CLINICAL TRIALS REPORTING PROGRAM (CTRP) SYSTEM

Warning Notice

- This warning banner provides privacy and security notices consistent with applicable federal laws, directives, and other federal guidance for accessing this Government system, which includes (1) this computer network, (2) all computers connected to this network, and (3) all devices and storage media attached to this network or to a computer on this network.
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NOTIFICATION TO RESPONDENT OF ESTIMATED BURDEN

OMB#: 0925-0600 EXP. DATE: 10/31/22

Public reporting burden for this collection of information is estimated to average sixty (60) minutes for this questionnaire, including the time to review instructions, search existing data sources, gather and maintain the data needed, and complete and review 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 current, valid OMB control number. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing the burden to NIH, Project Clearance Branch, 6705 Rockledge Drive, MSC 7974, Bethesda, MD 20892-7974, ATTN: PRA (0925-0600).

Do not return the completed form to this address.



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Per NCI	CTRP Regi	stration					
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National Cancer Institute		
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Amendment Trial Use this form to register trials with the NCI Clinical Trials Reporting Program. Required fields are marked by asterisks (*). Collapse All Collapse All		
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Lead Organization Trial Identifier:*	test 123	0
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Other Identifiers*		
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Lead Organization:*	Dana-Farber Cancer Institute	
Principal Investigator:*	Izbicki, Michael	Look Up Person
	Izbicki, Michael	Look Up Person
ponsor/Responsible Party	Izbicki, Michael	Look Up Person
ponsor/Responsible Party information in this section is REQUIRED to enable "Upload from NCI CTRP" in ClinicalTrials.gov	Izbicki, Michael Dana-Farber Cancer Institute	Look Up Person
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NIH Grant Information (for NIH funded Trials)*						×	
To record grant information, provide values for all fields, and then click the Add Grant button.							
	Is this trial funded by an NCI grant?	* • Yes O No					
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Existing Trial Related Documents						
Document Types	File Name					
IRB Approval Document	test-2.docx					
Change Memo Document	test - 3.docx					
Protocol Document	test-2.docx					
TSR	TSR_NCI-2017-00782_2017-07-21-1245_A123-rtf					
Trial Related Documents *	· · · · · · · · · · · · · · · · · · ·					
Amendment Related Documents To ensure successful registration, upload a Protocol document and an IRB Approval document. If the Protocol document does not include the CTRP accepts most standard document types. For additional information about what document types are accepted, please refer to the Help s	Informed Consent and/or participating sites, upload the Informed Consent document and a list of participating sites separately. You can use the Participating Sites template to submit your list of participating sites.					
Protocol Document.*	Browse					
Change Memo Document:**	Browse					
Protocol Highlighted Document:**	Browse					
IRB ApprovaL*	Browse					
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