

NCI CTRP Attachment 2C

NCI CTRP Accrual 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 clicks “Login”

Step 3: User enters “Email Address” and “Password” – see screenshot, page 3

Step 4: User reviews NCI Clinical Trials Reporting Program burden statement – see screenshot, page 4

Step 5: System displays “Search Submitted Clinical Trials” page – see screenshot, page 5

Step 6: User selects to “Submit Study Subject Accrual Information” and submits subject level accrual information on a registered trial – see screenshots, pages 6 - 9

CTRP Home page

NCI Registry - Mozilla Firefox

File Edit View History Bookmarks Tools Help

CTRP Login Screen

National Cancer Institute

U.S. National Institutes of Health | www.cancer.gov

NCI CTRP Registration Site

Log In

NCI CTRP

- Home
- Create Account
- Register Trial
- Search Trials
- Log In**
- Help

QUICK LINKS

- Clinical Trials Reporting Program (CTRP)
- Useful Templates and Documentation
- National Cancer Institute (NCI)
- NCI Center for Bioinformatics (NCICB)
- caBIG™ - Cancer Biomedical Informatics Grid™

Login

Help

Please log in to search, view and register clinical trial details. If you do not have an account, you may [create an account](#).





Email Address:

Password:

[\(Forgot Your Password?\)](#)
[\(Create an Account\)](#)

Log In

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CTRP Burden Statement

NCI CTRP

- Home
- My Account
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- Log Out
- Help

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NCI CLINICAL TRIALS REPORTING PROGRAM (CTRP) SYSTEM

OMB#: XXXX-XXXX EXP. DATE: XXXX/XXXX

NOTIFICATION TO RESPONDENT OF ESTIMATED BURDEN

Public reporting burden for this collection of information is estimated to average fifteen (15) 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.

Accept

Reject



Search Submitted Clinical Trials - Mozilla Firefox

File Edit View History Bookmarks Tools Help

https://trials-qa.nctn.org/

Search Submitted Clinical Trials

Most Visited Getting Started Latest Headlines

National Cancer Institute U.S. National Institute of Health

NCI CTRP Registration Site

Welcome, nshimko@scenpro.com

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Search Submitted Clinical Trials

Title:

Phase:

Identifier Type:

Organization Type:

Purpose:

Identifier:

Organization:

(e.g. NCI-2008-00015; ECOG-1234, etc)

Search My Trials: Search the trials I have submitted.
Search All Trials: Search all trials I have submitted as well as those registered by others.

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NATIONAL CANCER INSTITUTE | USA.gov

http://www.cancer.gov/ trials-qa.nci.nih.gov



Submit Study Subject Accrual Information



NCI CTRP

- Home
- Trial Search
- Submissions
- Study Subject Search
- Log Out

NCI-2009-00098: trial to test nullification err-Changed to generic contact for resp party

Lead Organization Trial ID: Trail to test nullify

Principal Investigator: Name, Lead

Lead Organization: Sponsortest - modified

Accrual Submissions

3 items found, displaying all items. 1

| Label | Description | Cut-off Date | Created Date | User Created | Submitted Date | User Submitted |
|--|------------------------------|--------------|--------------|---------------------------------|----------------|------------------------|
| Test Submission | This is a sample description | 09/18/2009 | 09/22/2009 | edmond.mulaire@semanticbits.com | 09/22/2009 | edmond.mulaire@semanti |
| Test Accrual Application | Initial testing | 09/09/2010 | 09/24/2009 | lanthone@gmail.com | 09/24/2009 | lanthone@gmail.com |
| Demo 1 | For accrual demo | 09/28/2009 | 09/28/2009 | nshimko@scenpro.com | | |

QUICK LINKS

- Clinical Trials Reporting Program (CTRP)
- National Cancer Institute (NCI)
- NCI Center for Bioinformatics (NCIB)
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NCI CTRP

- Home
- View Trials
- Search Trial
- Accrual Submission
 - View Previous Submission
 - New Submission

NCI-2009-00996: Phase II study for Melanoma Vaccine, or MDX-010/MDX-1379

Combination Treatment for Patients With Melanoma

Lead Organization Trial ID: S0919

Principal Investigator: Adjani, Anjali

Lead Organization: : Ochsner Health System

Participation Site: Ochsner Cancer Institute

New Accrual Submission

[View Patients](#)

QUICK LINKS

- National Cancer Institute (NCI)
- NCI Center for Bioinformatics (NCICB)
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Submission Title :

Submission Cut off Date :  (MM/YYYY)

Description:





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NCI-2009-00098: trial to test nullification err-Changed to generic contact for resp party

Lead Organization Trial ID: Trail to test nullify

Principal Investigator: Name, Lead

Lead Organization: Sponsortest - modified

Search Study Subject

Study Subject ID

Participating Site --Select--

Study Subject Birth Date (MM/YYYY):

Record Status: --Select--

List of Study Subjects

Nothing found to display.

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Lead Organization Trial ID: Trail to test nullify

Principal Investigator: Name, Lead

Lead Organization: Sponsortest - modified

Add Study Subject

Study Subject ID: *

Study Subject Birth Date (MM/YYYY): *

Study Subject Gender: * --Select-

Study Subject Race: * --Select-

Study Subject Ethnicity: * --Select-

Study Subject Country: * -Select-

Study Subject Zip Code:

Registration Date (mm/dd/yyyy)

Study Subject method of payment: --Select-

Disease: *

Participating Site * --Select-