

## **NCI CLINICAL TRIALS REPORTI**

### **NOTIFICATION TO RESPOND**

OMB#: 0925-0600 EX

Public reporting burden for this collection of information is es  
in combination with initial trial registration, including the time  
and maintain the data needed, and complete and review the c  
sponsor, and a person is not required to respond to, a collecti  
control number. Send comments regarding this burden estim  
including suggestions for reducing the burden to NIH, Project  
Bethesda, MD 20892-7974, ATTN: PRA (0925-0600) Do not ret

## **ING PROGRAM (CTRP) SYSTEM**

### **ENT OF ESTIMATED BURDEN**

(P. DATE: 02/28/2026

Estimated to average sixty (60) minutes for this questionnaire  
to review instructions, search existing data sources, gather  
collection of information. An agency may not conduct or  
ion of information unless it displays a current, valid OMB  
ate or any other aspect of this collection of information,  
Clearance Branch, 6705 Rockledge Drive, MSC 7974,  
urn the completed form to this address.

# **CTRP Trial Registration Participating Site Specification for C**

The participating sites template is designed for recording participating site data for interven

## **About this Document**

This document provides you with everything you need to upload clinical trial participating si

### **Template Instructions**

The Template Instructions worksheet provides detailed instructions for preparing y

### **Participating Site Data Specification and Collaborator Data Specification**

The specifications worksheets includes the following information:

- 1 Data elements
- 2 Order in which the data elements *must be* presented
- 3 Data element requirements
- 4 Valid values. The system accepts only those values listed in this documer
- 5 Notes. Additional information that helps you to ensure successful submis

### **Participating Site and Collaborator Examples**

These worksheets provide examples of a typical participating sites/collaborator dat

## Complete Trials

tional trials, especially if site-specific data is not included in the trial protocol.

tes and collaborator data to the CTRP Trial Registration system, including the following:

our data and uploading them to the system.

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sion of your data

a file.

# How to S1

## Main Step

- 1 Prep
- 2 Uplc

## Preparing

1 Ens

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- \*

2 Prep

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## Uploading

Upload your f

For detailed in

<https>

# Submit Complete Trial Participating Sites Data to the CTRP Trial Registration Application

## Steps for Uploading Your Data

Prepare the trial data file  
Upload the file in the Trial-Related Documents section in the Registration application

### 3. Trial Data Files

Ensure that your trial conforms to the supported criteria. This template supports the following:

Interventional trials

Complete trials (Data Table 4 Funding Sponsor Category is National, Externally Peer Reviewed, or Institutional)

Processing Statuses for trial updates: Accepted and beyond

Processing Statuses for trial amendments: Abstraction Verified Response or Abstraction Verified No Response

Prepare an Excel spreadsheet (.xls) containing the mandatory and optional data for the trial(s) as specified in the template.

**You must adhere to the following requirements:**

Enter trial elements required for registration in the order specified in the Participating Site Data Spec tab in this application.

Do not change the spelling of data elements or valid values.

Conform to the valid values guidelines when entering trial data.

Identify each trial uniquely

Participating site information must include the following data elements:

\* Study participating site data

\* At least one study site investigator's information

\* Participating site primary or central contact information. Generic contact information is accepted.

\* Organization attribute

\* Current recruitment status

\* Status date

\* Target accrual. This is mandatory if the target accrual is for a study at a participating site or if the lead or primary study site investigator's information must include the following data elements:

Study site investigator's information must include the following data elements:

\* Study site investigator data with person's attributes

\* Investigator's role in the study at the site.

See the [Participating Site Data Spec](#) for more information.

Participating site contact information is optional if the contact person is the investigator, or if the central contact person is the investigator.

Identify all persons and organizations with PO-IDs.

Note: You can request a list of CTRP persons and organizations along with PO-IDs from the CTRO at [nci.ctrp@nih.gov](mailto:nci.ctrp@nih.gov).

Or, you can use the organization/person lookup features in the CTRP Trial Registration application to search for persons and organizations.

Note: Although you can update Program Codes via the NCI CTRP Registration site, you can not update them via the

## **g Your File**

file in the Trial-Related Documents section of the CTRP Registration Site's Register Trial page.

Instructions for registering trials, refer to the NCI CTRP Reporting Program Registration Site User's Guide at <https://wiki.nci.nih.gov/x/7ZF4B>

r Center. Optionally, provide a local trial identifier.

Element order	Element	Required?
	<b>Study participating site data</b>	
1	Site #	Yes
2	Local Trial Identifier	
3	[Site] Organization PO-ID	
4	[Site] Name	Yes if PO-ID is not provided
5	[Site] Street Address	Yes if PO-ID is not provided
6	[Site] City	Yes if PO-ID is not provided
7	[Site] State/Province (US/Canada/Australia)	Yes for US, Canada, Australia and if PO-ID is not provided
8	[Site] Zip/Postal code	Yes if PO-ID is not provided
9	[Site] Country	Yes if PO-ID is not provided
10	[Site] Email	Yes if PO-ID is not provided
11	[Site] Phone	
12	[Site] Phone extension	
13	[Site] TTY	
14	[Site] FAX	
15	[Site] URL	
16	Is it NCI designated cancer center?	Yes
17	Study Current Recruitment Status at site	Yes
18	Study Current Recruitment Status date	Yes
19	Site Target Accrual	Yes if either site or lead organization is cancer center
20	Program Code	
	<b>Study site investigator's information</b>	
21	Investigator's Person PO-ID	
22	Investigator's First Name	Yes if PO-ID is not provided
23	Investigator's Middle name	
24	Investigator's Last Name	Yes if PO-ID is not provided
25	Investigator's Email	Yes if PO-ID is not provided
26	Investigator's Phone	Yes if PO-ID is not provided
27	Investigator's Phone Extension	

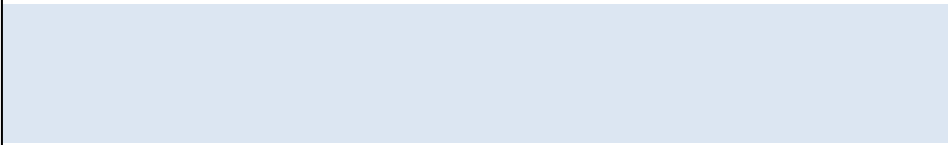
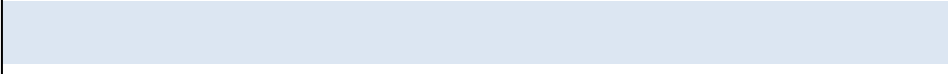
28	Investigator's Street Address	Yes if PO-ID is not provided
29	Investigator's Zip/Postal Code	Yes if PO-ID is not provided
30	Investigator's City	Yes if PO-ID is not provided
31	Investigator's State/Province (US/Canada, Australia)	Yes if country is US, Canada, Australia and PO-ID is not provided
32	Investigator's Country	Yes if PO-ID is not provided
33	Investigator's TTY	
34	Investigator's FAX	
35	Investigator's URL	
36	Investigator's Role in the study	Yes
37	Use investigator as site contact for the study	Yes
	<b>Study/Site Contact information</b>	
38	Contact type	Yes
	Generic Contact	
39	Title for generic contact	Yes if generic contact is used
40	Contact Email	Yes if generic contact is used
41	Contact Phone	Yes if generic contact is used
42	Contact Phone Extension	
	Personal Contact	
43	Contact Person's PO-ID	
44	Contact Person's First Name	Yes if personal contact is used and PO-ID is not provided
45	Contact Person's Middle Name	
46	Contact Person's Last Name	Yes if personal contact is used and PO-ID is not provided
47	Contact Email	Yes if personal contact is used and PO-ID is not provided
48	Contact Phone	Yes if personal contact is used and PO-ID is not provided
49	Contact Phone Extension	
50	Contact Person's Street Address	Yes if personal contact is used and PO-ID is not provided

51	Contact Person's Zip/Postal Code	Yes if personal contact is used and PO-ID is not provided
52	Contact Person's City	Yes if personal contact is used and PO-ID is not provided
53	Contact Person's State/Province (US/Canada/Australia)	Yes for personal contact and if country is US, Canada, Australia
54	Contact Person's Country	Yes if personal contact is used and PO-ID is not provided
55	Contact Person's TTY	
56	Contact Person's FAX	
57	Contact Person's URL	

Allowed values	Note
	<b>For participating sites only; lead organization must be included if it is also a participating site</b>
	Order in the list of participating sites
	Trial identifier at site
	PO-ID or organization mandatory attributes is required
	US, Canada, Australia only for the case when PO-ID is not provided
	If exists and phone is provided
Yes/No	Indicate if organization is a NCI designated cancer center
Not yet recruiting; Recruiting; Enrolling by invitation; Active, not recruiting; Completed; Suspended; Terminated; Withdrawn	
Date in the format mm/dd/yyyy	Date that corresponds to the current recruitment status change
Number	Mandatory if either site or lead organization is cancer center
	Site-specific Data Table 4 program code for NCI designated cancer center
	<b>Several records per one participating site are accepted</b>
	Person PO-ID or all mandatory person attributes is required
	Only initials
	Email address specific to study
	Phone specific to study
	Mandatory if exists and PO-ID is not provided

	Only for US/Canada/Australia in case if PO-ID is not provided
Principal Investigator, Sub-Investigator	
Yes/No	IF YES is selected, investigator will play participating site contact role for the study and no other participating site contact will be required
Site-Specific, Study-specific or central	Provide single contact for the study (study-specific) or site-specific contact for each participating site. This attribute is not required if site's investigator is assigned as site contact. There is no need to replicate central contact in each participating site record if central contact is selected and provided in the first record
	Generic contact or personal contact is required
	Several records per one participating site are accepted in case of site-specific contact type
	Email address specific to study
	Phone specific to study
	Mandatory if exists
	PO-ID or all mandatory person attributes is required
	Several records per one participating site are accepted in case of site-specific contact type
	Only initials
	Email address specific to study
	Phone specific to study
	Mandatory if exists and the phone is provided

	Only for US/Canada/Australia in case if PO-ID is not provided







#	Element	Mandatory?	Value
	<b>Collaborator information</b>	<b>Optional</b>	
1	Collaborator #		
3	Collaborator Organization PO-ID		
2	Collaborator Name	Yes if PO-ID is not provided	
4	Collaborator Street Address	Yes if PO-ID is not provided	
5	Collaborator City	Yes if PO-ID is not provided	
6	Collaborator State/Province (US/Canada/Australia)	Yes for US, Canada, Australia and PO- ID is not provided	
7	Collaborator Zip/Postal code	Yes if PO-ID is not provided	
8	Collaborator Country	Yes if PO-ID is not provided	
9	Collaborator Email	Yes if PO-ID is not provided	
10	Collaborator Phone		
11	Collaborator Phone extension		
12	Collaborator TTY		
13	Collaborator FAX		
14	Collaborator URL		
15	Collaborator role on the study	Yes if PO-ID is not provided	Funding Source, Agent Source, Laboratory

Note	
Order in the list of collaborators	
PO-ID or all mandatory organization attributes is required	
US, Canada, Australia only in case if PO-ID is not provided	
If exists and phone is provided	

	1	2	3	4	5
Site info	Site #	Local Trial Identifier	[Site] Organization PO-ID	[Site] Name	[Site] Street Address
<b>Study 1</b>					
	1	LI01		The University of Texas M. D. Anderson Cancer Center	1515 Holcombe Blvd
	1				
	2	LI02		<a href="#">Memorial Sloan-Kettering Cancer C</a>	1275 York Avenue
<b>Study 2</b>					
	1	LI04		<a href="#">Memorial Sloan-Kettering Cancer C</a>	1275 York Avenue
	2	LI06		The University of Texas M. D. Anderson Cancer Center	1515 Holcombe Blvd

6	7	8	9	10	11
[Site] City	[Site] State/Province (US/Canada)	[Site] Zip/Postal code (US/Canada)	[Site] Country	[Site] Email	[Site] Phone
Houston	TX	77030	USA	<a href="mailto:mailus@mdanderson.org">mailus@mdanderson.org</a>	(713) 792-5410
New York	NY	10065	USA	<a href="mailto:info@mskcc.org">info@mskcc.org</a>	212-639-2000
New York	NY	10065	USA	<a href="mailto:info@mskcc.org">info@mskcc.org</a>	212-639-2000
Houston	TX	77030	USA	<a href="mailto:mailus@mdanderson.org">mailus@mdanderson.org</a>	(713) 792-5410

12	13	14	15	16	17	18	19
[Site] Phone extension	[Site] TTY	[Site] FAX	[Site] URL	Is it NCI designated cancer center?	Study Current Recruitment Status at site	Study Current Recruitment Status date	Site Target Accrual
			<a href="http://www.mdanderson.org">http://www.mdanderson.org</a>	YES	recruiting	10/20/2008	55
			<a href="http://www.mskcc.org">http://www.mskcc.org</a>	YES	recruiting	11/2/2008	125
			<a href="http://www.mskcc.org">http://www.mskcc.org</a>	YES	recruiting	11/2/2008	125
			<a href="http://www.mdanderson.org">http://www.mdanderson.org</a>	YES	recruiting	10/20/2008	55

20		21	22	23	24	25
Program Code	Site Investigator	Investigator's Person PO-ID	Investigator's First Name	Investigator's Middle name	Investigator's Last Name	Investigator's Email
BM3			Mary	A	Simpson	<a href="mailto:msimpson@mdanderson.org">msimpson@mdanderson.org</a>
			Brandy	S	White	<a href="mailto:bwhite@mdanderson.org">bwhite@mdanderson.org</a>
			Helen	T	Harold	<a href="mailto:hharold@mskcc.org">hharold@mskcc.org</a>
			Helen	T	Harold	<a href="mailto:hharold@mskcc.org">hharold@mskcc.org</a>
BM3			Mary	A	Simpson	<a href="mailto:msimpson@mdanderson.org">msimpson@mdanderson.org</a>

26	27	28	29	30	31	32
Investigator's Phone	Investigator's Phone Extension	Investigator's Street Address	Investigator's Zip/Postal Code (US/Canada)	Investigator's City	Investigator's State/Province (US/Canada)	Investigator's Country
(713) 792-5410	235	1515 Holcombe Blvd	77030	Houston	TX	USA
(713) 792-5410	254	1515 Holcombe Blvd	77030	Houston	TX	USA
212-639-2000	3224	1275 York Avenue	10065	New York	NY	USA
212-639-2000-145	3224	1275 York Avenue	10065	New York	NY	USA
(713) 792-5410	235	1515 Holcombe Blvd	77030	Houston	TX	USA

33	34	35	36	37		38	
Investigator's TTY	Investigator's FAX	Investigator's URL	Investigator's Role in the study	Use investigator as site contact for the study	Site Contact Info	Contact type	Genetic Contact
			Principal Investigator	YES		Site-Specific	
			Sub-investigator	NO			
			Principal Investigator	NO		Site-Specific	
			Principal Investigator	NO		Study_specific	
			Principal Investigator	NO			





56	57		
Contact Person's FAX	Contact Person's URL		<b>Note</b>
			Site is a NCI designated cancer center, includes 2 investigators. One of the investigators is selected as this site contact.
			Site is a NCI designated cancer center, includes 1 investigator. Site-Specific contact is used (investigator is not used for site contact)
			Generic study-specific contact is used; no need to provide contact for each site separately.

1	3	2	4	5	6
Collaborator #	Collaborator Organization PO-ID	Collaborator Name	Collaborator Street Address	Collaborator City	Collaborator State/Province (US/Canada)
1		Publitek, Inc. dba Fotosearch	21155 Watertown Road	Waukesha	WI
2		<a href="#">AstraZeneca International</a>	P.O. Box 15437	Wilmington	DE

7	8	9	10	11	12
Collaborator Zip/Postal code (US/Canada)	Collaborator Country	Collaborator Email	Collaborator Phone	Collaborator Phone extension	Collaborator TTY
53186-1898	USA	<a href="mailto:mailus@fotosearch.com">mailus@fotosearch.com</a>	1-800-827-3920		
19850-5437	UK	<a href="mailto:info@astrazeneca-us.com">info@astrazeneca-us.com</a>	1 302 886 3000		

13	14	15
Collaborator FAX	Collaborator URL	Collaborator role on the study
262-717-0745	<a href="http://www.fotosearch.com">http://www.fotosearch.com</a>	Laboratory
302 886 2972	<a href="http://www.astrazeneca-us.com">www.astrazeneca-us.com</a>	Agent Source