Recent changes to CTRP 20210625

This page provides an overview of the changes in CTRP. For any questions or issues regarding these feature enhancements, please contact the CTRO (N CICTRO@mail.nih.gov).

June 2021

Transition to the NCIs multi-factor authentication system, Okta: Your CTRP username is now your institutional email address.

Migration of CTRP user accounts to Okta. Okta multi-factor authentication is used to provide additional security beyond just entering a username and password. Each Okta account requires the configuration of at least one of the following methods:

- Okta Verify: Use a 'Push Notification' sent to the Okta app on the users mobile device.
- SMS Authentication: Enter a single-use code sent to the users mobile device.
- Voice Call Authentication Phone call authentication by following voice instructions.

New CTRP login page

Clinical	l Trials Reporting Program
NIF	NIH Login
	OR
Username	
Password	
Remember	me
	SIGN IN
Need help signi	ing in?
Don't have an	account? Sign up

For additional information about creating and setting up an Okta account, please visit these CTRP User Guide pages:

Creating a CTRP Account

CTRP Okta Account Setup

March 2021

Updates to the Data Table 4 (DT4) Anatomic Site values displayed on the CTRP-generated DT4 report:

Previous DT4 Anatomic Site Value	Updated DT4 Anatomic Site Value
Leukemia, not otherwise specified	Leukemia, other
Breast – Female	Breast
Breast – Male	Breast
Hodgkin's Lymphoma	Hodgkin Lymphoma
Non-Hodgkin's Lymphoma	Non-Hodgkin Lymphoma
Soft Tissue / Sarcoma	Soft Tissue

Note: For trials with numerous "Primary Sites" (DT4 Anatomic Sites), the value "Multiple" for the DT4 Anatomic Site Code will be used and displayed on the CTRP-generated DT4 report. All existing trials in CTRP have been updated to reflect this change as well.

For a complete list of DT4 anatomic site values displayed on the CTRP-generated DT4 report, please visit this CTRP User Guide page:

https://wiki.nci.nih.gov/display/CTRPdoc/Data+Table+4+Anatomic+Site+Values

Removal of the following fields from the Trial Summary Report (TSR):

- Keywords
- Program Code in the Data Table 4 Section
 Target Accrual in the Participating Sites Section
- Reporting Data Set Method

September 2020

CTRP Trial Record Verification: Improved the user experience for CTRP trial owners, trial submitters and site administrators with Verifying, Updating and /or Amending their trial records. Key highlights from this enhancement include:

1. New page highlighting all Trials Needing Verification

Clinica	I Trials Repor	The Trials Nee approaching th Verification D	eir Record V	erification	Due Date	or the Record	d	ctrpsubstractor	r Cl 👻
🗹 Trials to Veril	y ₉₀ Q Search	verify their trial						? Help	
rials Needin	g Verification								
he trials below are either ial information is accurate	approaching their Record Verifica e and up to date.	ation Due Date or the Record	Verification Due Date is in	h the past. The NCI	requests that trial ov	wners verify their trial reco	rds in CTRP at least every 6	i months to ensure that	the:
how 10 🗸				Search:		Choose columns <<	< 1 2 3 4	59>	>>
NCI Trial Identifier 🖣	Title	ne NCI Trial Ide e trial record.	ntifier to	n Date 🔺 Lead (Organization \$	Lead Org Trial Identifier	Available Actions	Current Trial State	us Ø
NCI-2017-00384	A Phase I/II S With Multi-Ag			Case We Univers		PCC 348-02	Select Action 👻	In Review	1
NCI-2017-00389	A Phase I/II Study Of Brentuxim With Multi-Agent Chemotherap		04/11/2017	Case We Univers		PCC 34890534-4	Select Action 👻	In Review	1
NCI-2017-00385	A Phase I/II Study Of Brentuxim With Multi-Agent Chemotherap		04/11/2017	Case We Univers		PCC 348-02 II	Select Action 👻	In Review	1
NCI-2017-00381	A Phase I/II Study Of Brentuxim With Multi-Agent Chemotherap		04/11/2017	Case W Univers		PCC 348-02h	Select Action 👻	In Review	1
NCI-2017-02638	New Test Trial		06/28/2017	Case Co Cancer		PCC 86422673 ss410	Select Action 👻	In Review	,
NCI-2017-02636	test73834		06/28/2017	Case Co Cancer		est783484	Select Action 👻	In Review	

2. Ability to view all trial details on the Verify screen

Update/Verify Trial I Trial Details	A Phase I/II Study Of Bxim			Trial Identifiers			Other A	tions
				Total I deve ald a se			will allow	users
	combination martiners	ab Vedotin In gent Chemotherapy		NCI Trial Identifier	NCI-2017-00331		Amend a when up the Prote	dating
Trial owners, trial subr review and edit trial da Update/Verify button.	ata prior to sele	cting the		Identifier*	UPCC 34890534-434 13 characters left NCT01994552	0	IRB App This is a available	roval so
Verification Method, the Trial Data Verifica Trial Data page. The p	ation section o	n the Update/Verify	,	Other Identifiers			Trials No Verificat and Tria	eedin
Lead Organizatio	on / Principal Investi	gator		Regulatory Informat	tion	Amend Change Status	Search under Se Actions	elect
Lead Organization	National Cancer Institute Prevention			The information in this section ClinicalTrials.gov Studies a U.S. FDA-regula	on is REQUIRED to enat	View TSR View XML View Details	\square	

- 3. Simplified workflow allowing users to Verify, Update or Amend trials within same form
- 4. New monthly email notification for alerting trial owners, trial submitters,
- and site administrators of Trials Needing Verification

Date: 07/31/2020 Dear firstname lastname, The following trial(s) are due fo	or Data Verification in CTRP:		Users will now receive monthly email notifications regarding trials due and past due for Data Verification. For example, trials due for Data Verification in May will start receiving the				
NCI ID NCI-2017-02629	Lead Organization Trial ID test7382	Verification Due Date 08/20/2020	email notification in April. The notification will continue monthly until all trials due for Data Verification, both present and past, have been verified				
The following trial(s) are past d	ue for Data Verification in CTRP:						
NCI ID NCI-2017-02619 NCI-2017-02623 NCI-2017-02625	Lead Organization Trial ID test945798 test237823 test648843	Verification Due Date 06/28/2018 06/28/2018 07/07/2018					
 Date the last data abstractive Date the last Update was subsequence Date the last Data Verification 	on was verified (both new registra ubmitted in CTRP tion was performed in CTRP		fication due date is determined based on the following: ssions) in CTRP				
 Select the trial you wish to If the data is accurate and 		d y button	w/registry/protected/view.VerifyTrials.action				
If you have questions on this pro	ocess, please contact us at ncictro	@mail.nih.gov					
Thank you for your participation	n in the NCI Clinical Trials Report	ting Program (CTRP).					

For more information on the CTRP Trial Record Verification processes, please visit these CTRP User Guide pages:

https://wiki.nci.nih.gov/display/CTRPdoc/Updating+Trial+Information

https://wiki.nci.nih.gov/display/CTRPdoc/How+to+Verify+Trial+Data

https://wiki.nci.nih.gov/display/CTRPdoc/How+to+Access+the+Trial+Data+Verification+Page

https://wiki.nci.nih.gov/display/CTRPdoc/Amending+Trials

July 2020

<u>Flexible Accrual</u>: Enhanced the CTRP accrual reporting capabilities to include more "Flexible Accrual" reporting options. Centers can now request to update the default accrual reporting type on a study registered in CTRP. Changing the default accrual reporting type requires a ticket to be submitted to the CTRO (NCICTRO@mail.nih.gov) for review and approval, before any change is granted.

1. The current default accrual reporting types are as follows:

Trial Type	Default Accrual Reporting Type
Complete Interventional	Subject
Complete Non-Interventional	Subject
Abbreviated/Imported Interventional	Summary
Abbreviated/Imported Non-Interventional	Summary

 CTRP is also introducing a new accrual reporting type, "Partial Subject." Partial Subject accrual requires the reporting of Study Subject ID, Registration Date and Participating Site data only. The other accrual data fields are optional. The reporting of Partial Subject accrual also require s a ticket to be submitted to the CTRO (NCICTRO@mail.nih.gov) for review and approval, before any change is granted.

3. Accrual Data Element changes:

- a. <u>Study Subject Birth Date</u>: For any/all ways to report accrual, only the month and year (MM/YYYY) are required to be collected.
 <u>Study Subject Country</u>: Required for all methods of reporting accrual (was previously optional)
- c. <u>Disease</u>:
 - i. Study Subject: Required
 - ii. Partial Subject: Not required. However, for ICD-O-3 Disease Codes if Disease is reported, then Site must be reported.
- d. <u>Site</u>:
 - i. Study Subject: Required
 - ii. Partial Subject: Not required. However, for ICD-O-3 Disease Codes if Site is reported, then Disease must be reported.
- e. Disease Codes: CTRP uses a hybrid model with the codes from IACR and the codes in NCI SEER that are not available in IACR.

For more information on the CTRP Accrual process, including Flexible Accrual, Partial Subject accrual and other changes made within the Accrual application (e.g., ability to Delete All Trial Study Subjects,) please visit these CTRP User Guide pages:

https://wiki.nci.nih.gov/display/CTRPdoc/About+Accrual

https://wiki.nci.nih.gov/display/CTRPdoc/Searching+for+Trials+in+Accrual

If you would like to receive future CTRP announcements you may add yourself to the list here: https://list.nih.gov/cgi-bin/wa.exe?SUBED1=ctrp-users-l&A=1