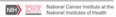
OMB #0925-xxxx Expiration Date: xx/xx/xxxx



Team Driven. Cancer Therapy Focused.

Experimental Therapeutics Clinical Trials Network



Introduction

The National Cancer Institute (NCI) would like to know about your experiences with the Experimental Therapeutics Clinical Trials Network (ETCTN) over the past grant year (April 2014 - March 2015).

Your input will help NCI assess and refine the ETCTN processes and identify areas for improvement.

The survey will ask questions about ETCTN processes and activities and trial portfolio. It should take approximately 15 minutes to complete.

Your responses are confidential and all results will be reported in the aggregate. Your participation and responses will have no bearing on your ETCTN grant, or any future interactions with NCI.

We thank you for your assistance!

To continue and begin the survey, click the "Next" button below.

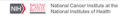


If you experience any technical difficulties, please contact the survey administrator, User-Centered Design at survey@user-centereddesign.com



Team Driven. Cancer Therapy Focused.

Experimental Therapeutics Clinical Trials Network



OMB# 0925-0046-09 Exp. Date: 05/31/2016

Privacy Statement and Consent

Collection of this information is authorized by The Public Health Service Act, Section 411 (42 USC 285a). Participation is voluntary and there are no penalties for not participating or withdrawing from the study at any time. The information collected in this study will be kept private to the extent provided by law. Names and other identifiers will not appear in any report of the study. Information provided will be combined for all study participants and reported as summaries. You are being contacted by User-Centered Design, Inc. via email to complete this instrument so that we can gain feedback from you on your experiences and recommendations regarding the implementation of the ETCTN program.

Please click the "Next" button if you consent to taking this survey.

Public reporting burden for this collection of information is estimated to average 15 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 208927974, ATTN: PRA (0925-0046-09). Do not return the completed form to this address.



perimental Therapeutics Clinical Trials Network	NIH	National Cancer Institute National Institutes of He
	_	Opt-out of sur
	•	5% Comp
Role and Involvement with ETCTN		
The first set of questions asks about your role within the ETCTN and your overall experier first grant year (April 2014 - March 2015).	nces with the pro	gram during th
Please review the following ETCTN staff roles and descriptions and select the primary roctivities for the program. If other roles describe your activities to a lesser degree, please se		
, , , , , , , , , , , , , , , , , , , ,		
		Secondary
	Primary role (select one)	role (select all that apply)
Grant administrator: Primary administrator for the program; makes roster changes in RSS; primary contact for program matters, including study announcements; disseminates ETCTN information from NCI to sites.		(select all
RSS; primary contact for program matters, including study announcements; disseminates	(select one)	(select all that apply)
RSS; primary contact for program matters, including study announcements; disseminates ETCTN information from NCI to sites. Site administrator: Primary contact person at each LAO or P2C-participating site; makes oster updates for their sites via CTSU website; disseminates ETCTN information to site	(select one)	(select all that apply)
RSS; primary contact for program matters, including study announcements; disseminates ETCTN information from NCI to sites. Site administrator: Primary contact person at each LAO or P2C-participating site; makes oster updates for their sites via CTSU website; disseminates ETCTN information to site staff. Study administrator: Primary contact person at a LAO or P2C-participating site for a	(select one)	(select all that apply)
RSS; primary contact for program matters, including study announcements; disseminates ETCTN information from NCI to sites. Site administrator: Primary contact person at each LAO or P2C-participating site; makes oster updates for their sites via CTSU website; disseminates ETCTN information to site taff. Study administrator: Primary contact person at a LAO or P2C-participating site for a particular ETCTN study.	(select one)	(select all that apply)

	Team Dri			
xperimental Therapeutics Clinical Trials Network			NIH Na	itional Cancer Institute a itional Institutes of Heal
				Opt-out of surv
		_		10% Comple
Thinking about your experiences with the ETCTN this past grant year (Apne program?	ril 2014 - Mai	rch 2015), ho	ow involved	d were you
 Extremely involved Very involved Somewhat involved Slightly involved Not at all involved 				
. Think about your experiences with the ETCTN this past grant year and incelow.	Strongly disagree	uch you agn	ee with the	Strongly agree
At the beginning of the ETCTN, I had a very clear understanding of how the program operated.	0	0	0	0
currently have a very clear understanding of how the ETCTN operates.	0	0	\circ	0
have a clear understanding of my role.	0	0	0	0
< Back		S	ave and C	Continue >
Your information will be saved every time you hit "save and continue". You may close If you experience any technical difficulties, plea survey administrator, User-Centered Design at <a and="" any="" close="" continue".="" difficulties,="" experience="" href="mailto:survey@usersenger</th><th>se contact the</th><th>is survey anyti</th><th></th><th></th></tr><tr><td>Your information will be saved every time you hit " if="" may="" plea<="" save="" td="" technical="" you=""><td>se contact the ser-centereddes</td><td>is survey anyti</td><td>me until it is</td><td>submitted.</td>	se contact the ser-centereddes	is survey anyti	me until it is	submitted.
Your information will be saved every time you hit "save and continue". You may close If you experience any technical difficulties, plea survey administrator, User-Centered Design at survey@u	se contact the ser-centereddes	is survey anyti	me until it is	submitted.
Your information will be saved every time you hit "save and continue". You may close If you experience any technical difficulties, plea survey administrator, User-Centered Design at survey@u • ETCTN	se contact the ser-centereddes	is survey anyti	me until it is	submitted.
Your information will be saved every time you hit "save and continue". You may close If you experience any technical difficulties, plea survey administrator, User-Centered Design at survey@u • ETCTN	se contact the ser-centereddes	is survey anyti	me until it is	submitted. Dy Focuse Sonal Cancer Institute a
Your information will be saved every time you hit "save and continue". You may close If you experience any technical difficulties, plea survey administrator, User-Centered Design at survey@u The property of	se contact the ser-centereddes	is survey anyti	me until it is	submitted. Dy Focuse tonal Cancer Institute tonal Institutes of Healt Opt-out of surv
Your information will be saved every time you hit "save and continue". You may close If you experience any technical difficulties, plea survey administrator, User-Centered Design at survey@u • ETCTN	se contact the ser-centereddes	is survey anyti	me until it is	submitted. Dy Focuse tonal Cancer Institute tonal Institutes of Healt Opt-out of surv
Your information will be saved every time you hit "save and continue". You may close If you experience any technical difficulties, plea survey administrator, User-Centered Design at survey@u The property of	se contact the ser-centereddes	is survey anyti	me until it is	Submitted. Dy Focuse tional Cancer Institute of Heal Opt-out of Surv 15% Comple
Your information will be saved every time you hit "save and continue". You may close If you experience any technical difficulties, plea survey administrator, User-Centered Design at survey@u **PROPERTY OF THE PROPERTY OF	se contact the ser-centereddes	is survey anyti	er Therap	submitted. Dy Focuse tional Cancer Institute a tional Institutes of Healt Opt-out of surv 15% Comple
Your information will be saved every time you hit "save and continue". You may close If you experience any technical difficulties, plea survey administrator, User-Centered Design at survey@u **ETCTN **xperimental Therapeutics Clinical Trials Network **Experience and Satisfaction with ETCTN Processes The next set of questions asks about your experience with the process to digrant year (April 2014 - March 2015). In the past grant year, were you involved in the process to submit a protocom.	se contact the ser-centereddes	is survey anyti	er Therap	submitted. Dy Focuse tional Cancer Institute a tional Institutes of Healt Opt-out of surv 15% Comple

Your information will be saved every time you hit "save and continue". You may close and return to this survey anytime until it is submitted.

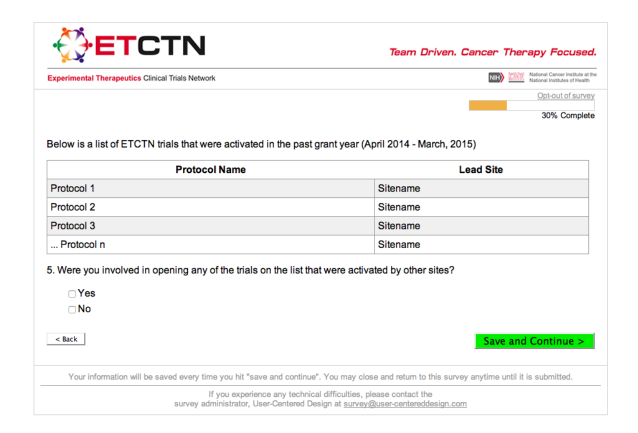
If you experience any technical difficulties, please contact the survey administrator, User-Centered Design at survey@user-centereddesign.com

Note: 4a appears conditionally if "yes" is selected in Q4.

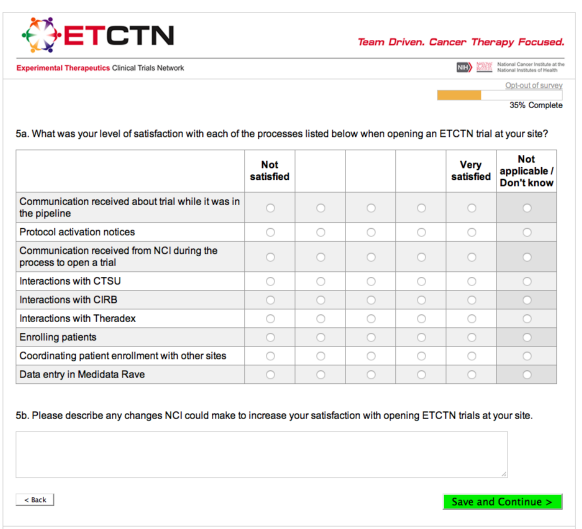
Experimental Therapeutics Clinical Trials Network		NIH	National Institutes of Hea
			Opt-out of sur
4a. Which activities did you participate in when submitting the protocol to NCI	at your site during th	e previous	20% Compl s grant year?
	Yes	No	Not applicable Don't know
Project Team Member Application (PTMA) submission	0	0	0
Letter of intent (LOI) submission	0	0	0
Protocol submission	0	0	0
Protocol approval	0	0	0
Informed consent document for CIRB-approved studies	0	0	0
CIRB approval	0	0	0
Protocol activation	0	0	0
Opening a protocol	0	0	0
Study build in Medidata Rave	0	0	0
Enrolling patients in a study	0	0	0
Coordinating patient enrollment with other sites / Reserving slots in OPEN	0	0	0
Study data entry into Medidata Rave	0	0	0
Study data entry into Medidata Rave < Back Your information will be saved every time you hit "save and continue". You may close an		Save an	d Contin

Note: 4b and 4c appear together conditionally if "yes" is selected for any activities in Q4a. 4b is populated with any activities identified as "yes" in Q4.

25% Complete the complete that was your level of satisfaction with the activities in which you participated? Not Very	Experimental Therapeutics Clinical Trials Network				NIH) NIH	lational Cancer Institute al lational Institutes of Health
b. What was your level of satisfaction with the activities in which you participated? Not satisfied Very satisfied Ver						Opt-out of surv
Not satisfied Project Team Member Application (PTMA) submission Letter of intent (LOI) submission Protocol submission Protocol approval Informed consent document for CIRB-approved studies CIRB approval Protocol activation Opening a protocol Study build in Medidata Rave Enrolling patients in a study Coordinating patient enrollment with other sites OR reserving slots in OPEN Study data entry into Medidata Rave c. Overall, what challenges did you encounter during these activities in the table above? (Select all that apply) Did not experience challenges Insufficient training Lack of helpdesk support			- 40			25% Comple
Project Team Member Application (PTMA) submission Letter of intent (LOI) submission Protocol submission Protocol approval Informed consent document for CIRB-approved studies CIRB approval Protocol activation Opening a protocol Study build in Medidata Rave Enrolling patients in a study Coordinating patient enrollment with other sites OR reserving slots in OPEN Study data entry into Medidata Rave c. Overall, what challenges did you encounter during these activities in the table above? (Select all that apply) Did not experience challenges Insufficient training Lack of helpdesk support	b. what was your level of satisfaction with the activities in which	Not	ea?			Very satisfied
Protocol submission Protocol approval Informed consent document for CIRB-approved studies CIRB approval Protocol activation Opening a protocol Study build in Medidata Rave Enrolling patients in a study Coordinating patient enrollment with other sites OR reserving slots in OPEN Study data entry into Medidata Rave C. Overall, what challenges did you encounter during these activities in the table above? (Select all that apply) Did not experience challenges Insufficient training Lack of helpdesk support	Project Team Member Application (PTMA) submission	0	0	0	0	
Protocol approval Informed consent document for CIRB-approved studies CIRB approval Protocol activation Opening a protocol Study build in Medidata Rave Enrolling patients in a study Coordinating patient enrollment with other sites OR reserving slots in OPEN Study data entry into Medidata Rave C. Overall, what challenges did you encounter during these activities in the table above? (Select all that apply) Did not experience challenges Insufficient training Lack of helpdesk support	Letter of intent (LOI) submission	0	0	0	0	0
Informed consent document for CIRB-approved studies CIRB approval Protocol activation Opening a protocol Study build in Medidata Rave Enrolling patients in a study Coordinating patient enrollment with other sites OR reserving slots in OPEN Study data entry into Medidata Rave c. Overall, what challenges did you encounter during these activities in the table above? (Select all that apply) Did not experience challenges Insufficient training Lack of helpdesk support	Protocol submission	0	0	0	0	0
CIRB approval Protocol activation Opening a protocol Study build in Medidata Rave Enrolling patients in a study Coordinating patient enrollment with other sites OR reserving slots in OPEN Study data entry into Medidata Rave Co. Overall, what challenges did you encounter during these activities in the table above? (Select all that apply) Did not experience challenges Insufficient training Lack of helpdesk support	Protocol approval	0	0	0	0	0
Protocol activation Opening a protocol Study build in Medidata Rave Enrolling patients in a study Coordinating patient enrollment with other sites OR reserving slots in OPEN Study data entry into Medidata Rave c. Overall, what challenges did you encounter during these activities in the table above? (Select all that apply) Did not experience challenges Insufficient training Lack of helpdesk support	Informed consent document for CIRB-approved studies	0	0	0	0	0
Opening a protocol Study build in Medidata Rave Enrolling patients in a study Coordinating patient enrollment with other sites OR reserving slots in OPEN Study data entry into Medidata Rave c. Overall, what challenges did you encounter during these activities in the table above? (Select all that apply) Did not experience challenges Insufficient training Lack of helpdesk support	CIRB approval	0	0	0	0	0
Study build in Medidata Rave Enrolling patients in a study Coordinating patient enrollment with other sites OR reserving slots in OPEN Study data entry into Medidata Rave C. Overall, what challenges did you encounter during these activities in the table above? (Select all that apply) Did not experience challenges Insufficient training Lack of helpdesk support	Protocol activation	0	0	0	0	0
Enrolling patients in a study Coordinating patient enrollment with other sites OR reserving slots in OPEN Study data entry into Medidata Rave Co. Overall, what challenges did you encounter during these activities in the table above? (Select all that apply) Did not experience challenges Insufficient training Lack of helpdesk support	Opening a protocol	0	0	0	0	0
Coordinating patient enrollment with other sites OR reserving slots in OPEN Study data entry into Medidata Rave C. Overall, what challenges did you encounter during these activities in the table above? (Select all that apply) Did not experience challenges Insufficient training Lack of helpdesk support	Study build in Medidata Rave	0	0	0	0	0
Study data entry into Medidata Rave c. Overall, what challenges did you encounter during these activities in the table above? (Select all that apply) Did not experience challenges Insufficient training Lack of helpdesk support	Enrolling patients in a study	0	0	0	0	0
c. Overall, what challenges did you encounter during these activities in the table above? (Select all that apply) Did not experience challenges Insufficient training Lack of helpdesk support		0	0	0	0	0
□ Did not experience challenges □ Insufficient training □ Lack of helpdesk support	Study data entry into Medidata Rave	0	0	0	0	0
☐ Unaware of other sites that have opened trial	Coordinating patient enrollment with other sites OR reserving slots in OPEN Study data entry into Medidata Rave c. Overall, what challenges did you encounter during these active Did not experience challenges Insufficient training Lack of helpdesk support Limited communication from NCI	0	0	0	0	0

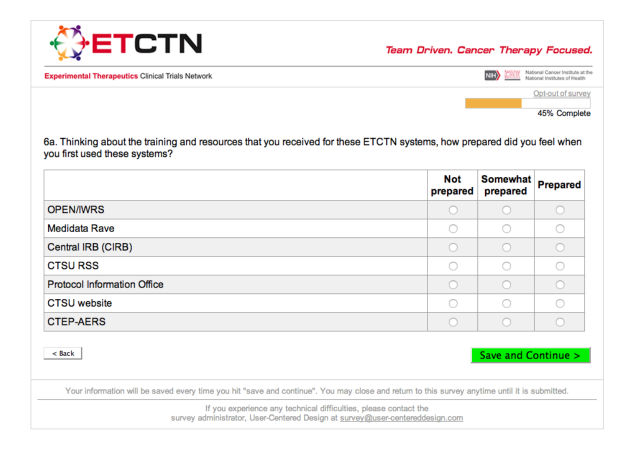


Note: 5a and 5b appear together conditionally; they appear if "yes" is selected in Q5.

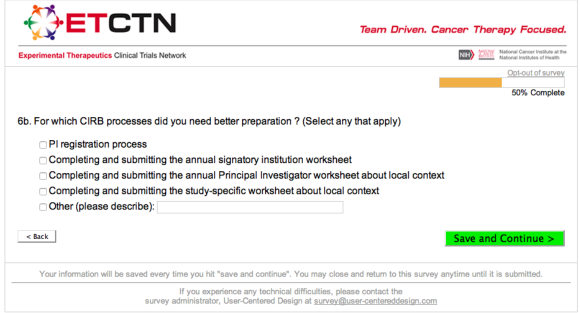


xperimental Therapeutics Clinical Trials Network				NIH	National Cancer Institut National Institutes of He
					Opt-out of su
					40% Comp
Experience and Satisfaction with E	TCTN Systems and	d Resources			
The following questions ask about you		on with various c	entralized ETC	TN systems/servi	ices for the
past grant year (April 2014 - March 20	15).				
. In the past grant year, how often did y	ou use each of the f	ollowing ETCTN	l systems?		
				More than 5	Have not
	1 time	2-3 times	4-5 times	times	used
OPEN/IWRS	0	0	0	0	0
A PLANE	0	0	0	0	0
Medidata Rave					
Central IRB (CIRB)	0	0	0	0	0
	0	0	0	0	0
Central IRB (CIRB)	0	_		_	0
Central IRB (CIRB) CTSU RSS (Rosters)	0 0	0	0	0	0
Central IRB (CIRB) CTSU RSS (Rosters) Protocol Information Office	0 0 0	0	0	0	0
Central IRB (CIRB) CTSU RSS (Rosters) Protocol Information Office CTSU website	0 0 0	0 0	0	0	0
Central IRB (CIRB) CTSU RSS (Rosters) Protocol Information Office CTSU website CTEP-AERS	0 0 0	0 0	0	0 0	0 0
Central IRB (CIRB) CTSU RSS (Rosters) Protocol Information Office CTSU website	0 0 0	0 0	0	0 0	o o o

Note: 6a appears conditionally populated with any systems used 1 or more times in Q6.



Note: 6b appears conditionally, if the user has selected "not prepared" or "somewhat prepared" for any items in Q6a.



Note: 6c and 6d appear together conditionally; 6c appears populated with any systems used 1 or more times in Q6. 6d appears if 6c appears.

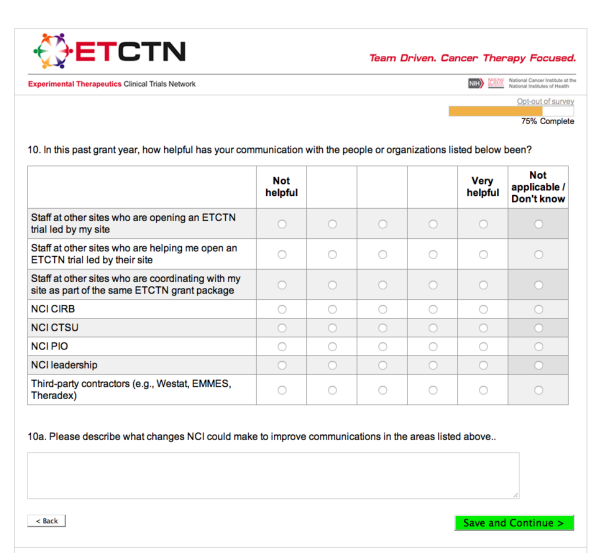
Experimental Therapeutics Clinical Trials Network				NIH)	National Cancer Institute National Institutes of Hea
					Opt-out of sur
					55% Compl
c. Overall, how satisfied were you with using the follow	owing ETCTN systems du	uring the pa	ast grant ye	ar?	
	Not satisfied				Very satisfied
OPEN/IWRS	0	0	0	0	0
Medidata Rave	0	0	0	0	0
Central IRB (CIRB)	0	0	0	0	0
CTSU RSS	0	0	0	0	0
D : 11 f :: 0ff		_	_	_	0
Protocol Information Office		0			
d. Looking at the list below, what could have helped our site? (check all that apply) Received training closer to utilizing ETCTN sy	you better prepare to use	0	ns required	0	0
CTSU website d. Looking at the list below, what could have helped our site? (check all that apply)	you better prepare to use	0		0	0
d. Looking at the list below, what could have helped our site? (check all that apply) Received training closer to utilizing ETCTN sy Helpdesk support Increased communication from NCI Increased communication with other sites con Shared "best practices" with other sites	you better prepare to use	0		0	0
d. Looking at the list below, what could have helped our site? (check all that apply) Received training closer to utilizing ETCTN sylone Helpdesk support Increased communication from NCI Increased communication with other sites con Shared "best practices" with other sites ETCTN Q&A documents CTSU Q&A documents ETCTN FAQ documents	you better prepare to use	the system		0	0
d. Looking at the list below, what could have helped our site? (check all that apply) Received training closer to utilizing ETCTN sylonger Helpdesk support Increased communication from NCI Increased communication with other sites con Shared "best practices" with other sites ETCTN Q&A documents CTSU Q&A documents ETCTN FAQ documents ETCTN process flowchart to describe initial did	you better prepare to use /stem ducting trial	the system		0	0
d. Looking at the list below, what could have helped our site? (check all that apply) Received training closer to utilizing ETCTN sylenger Helpdesk support Increased communication from NCI Increased communication with other sites con Shared "best practices" with other sites ETCTN Q&A documents CTSU Q&A documents ETCTN FAQ documents ETCTN process flowchart to describe initial decession of the site o	you better prepare to use /stem ducting trial	the system		0	0
CTSU website Sd. Looking at the list below, what could have helped your site? (check all that apply) Received training closer to utilizing ETCTN sylenger Helpdesk support Increased communication from NCI Increased communication with other sites con Shared "best practices" with other sites ETCTN Q&A documents CTSU Q&A documents ETCTN FAQ documents ETCTN process flowchart to describe initial difference in the country of th	you better prepare to use /stem ducting trial	the system		0	0
Helpdesk support Increased communication from NCI Increased communication with other sites con Shared "best practices" with other sites ETCTN Q&A documents CTSU Q&A documents ETCTN FAQ documents ETCTN process flowchart to describe initial describe initial describes to previously recorded ETCTN webing	you better prepare to use /stem ducting trial	the system	ns required	to open E	0

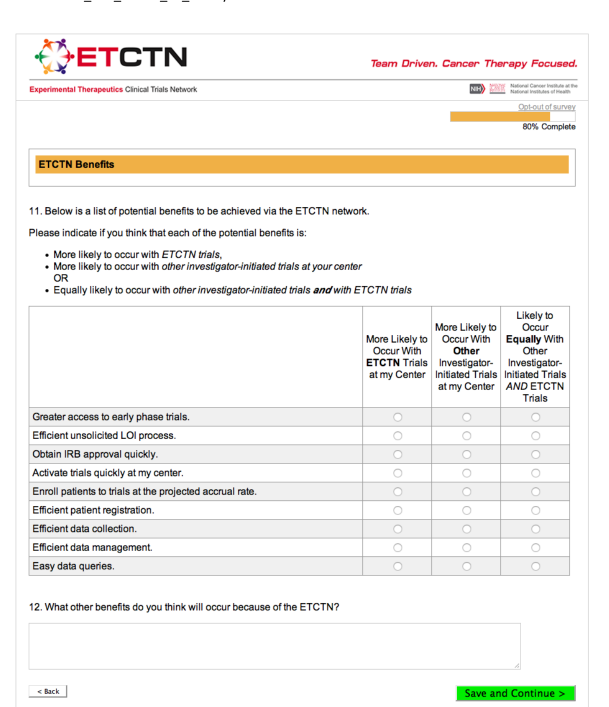
xperimental Therapeutics Clinical Trials Network				NIH Nat	tional Cancer Institute tional Institutes of Hea
					Opt-out of sun
The following questions ask about your use and satisfaction with year.	n various ce	entralized E	TCTN resou	rces for the	past grant
On average, how often do you use or refer to the following ETC	TN resourc	es?	T		T
	Daily	Weekly	Monthly	Quarterly or less	Have not used
TCTN Help Desks	0	0	0	0	0
CTSU bi-weekly broadcast	0	0	0	0	0
TCTN Trial update notifications	0	0	0	0	0
ETCTN training resources (e.g., checklists, information sheets, vebinars)	0	0	0	0	0
CTSU website	0	0	0	0	0
CIRB Helpdesk	0	0	0	0	0
IRB Helpdesk Are there additional resources that the ETCTN can provide to				0	0
				Save and C	ontinue >

Note: Q7b appears conditionally, populated with any resources used quarterly or more frequently in Q7.

b. How satisfied were you with using the following ETCTN resour	rces during th				Opt-out of sur
b. How satisfied were you with using the following ETCTN resour	rces during th				
b. How satisfied were you with using the following ETOTIV resour		na naet ara	nt vear?		00% Compi
	Not satisfied	ie past gra	int year:		Very satisfied
TCTN Help Desks	0	0	0	0	0
TSU bi-weekly broadcast	0	0	0	0	0
TCTN trial update notifications	0	0	0	0	0
TCTN training resources (e.g., checklists, information sheets, ebinars)	0	0	0	0	0
TSU website	0	0	0	0	0
IRB Helpdesk	0	0	0	0	0

ETCTN		Team Dri	ven. Canc	er Therap	oy Focused
Experimental Therapeutics Clinical Trials Network					tional Cancer Institute at the
					Opt-out of survey
					70% Complete
Experience and Satisfaction with ETCTN Communica	tion				
s. Looking at the items below, how well do you believe each ne past grant year?	n of the items below	v was comr	municated to	o your site t	oy NCI over
ETCTN goals	0	0	0	0	0
ETCTN protocols in development	0	0	0	0	0
Current ETCTN activated protocols	0	0	0	0	0
ETCTN trial updates	0	0	0	0	0
D. How do you typically obtain information about the ETCTN Trial-specific email updates directly from NCI CTSU's Bi-Weekly Broadcast email Communication from an administrator at your site Communication from an administrator at another site Early phase trial meetings at your site Grant PI at your site Other investigators at your site		ck all that a	ipply)		
CTSU website Other (please describe):					
□ CTSU website			2	save and C	ontinue >
CTSU website Other (please describe):					







Social Network Collaborations

We would like to learn more about your work with other researchers that are involved in the ETCTN.

13. Please look at the list of ETCTN organization members below and indicate a "yes" for those you had direct collaboration with over the past year (either developing protocols or opening trials).

ETCTN Lead Academic Organizations and Affiliates	Yes	No
Translational Genomics Research Institute	0	0
City of Hope Comprehensive Cancer Center	0	0
UC Davis Comprehensive Cancer Center	0	0
USC Norris Comprehensive Cancer Center	0	0
University of Colorado Cancer Center - Anschutz Cancer Pavilion	0	0
Yale Cancer Center	0	0
H. Lee Moffitt Cancer Center and Research Institute	0	0
Emory University/Winship Cancer Institute	0	0
University of Chicago	0	0
Johns Hopkins University/Sidney Kimmel Comprehensive Cancer Center	0	0
National Cancer Institute Developmental Therapeutics Clinic	0	0
University of Maryland Greenbaum Cancer Center	0	0
Dana-Farber Cancer Center	0	0
Massachusetts General Hospital	0	0
Wayne State University/Karmanos Cancer Institute	0	0
Mayo Clinic Rochester	0	0
Washington University	0	0
Rutgers University - Cancer Institute of New Jersey	0	0
Roswell Park Cancer Institute	0	0
Duke University	0	0
UNC Chapel Hill	0	0
Case Western Reserve University	0	0
Cleveland Clinic Foundation	0	0
Ohio State University Comprehensive Cancer Center	0	0
Fox Chase Cancer Center	0	0
University of Pittsburgh Cancer Institute	0	0
Vanderbilt-Ingram Cancer Center	0	0
University of Texas MD Anderson Cancer Center	0	0
Virginia Commonwealth University	0	0
University Wisconsin Carbone Cancer Center	0	0
British Columbia Cancer Agency	0	0
Juravinski Cancer Center	0	0
University Health Network/Princess Margaret Cancer Center	0	0

< Back Save and Continue >

Your information will be saved every time you hit "save and continue". You may close and return to this survey anytime until it is submitted.

Experimental Therapeutics Clinical Trials Network				NIH) Na Na	tional Cancer Institutional Institutes of H
					Opt-out of su
					90% Comp
Demographics					
The next set of questions asks about your background	I and experience with ca	ncer reseal	rch.		
Please indicate your sex:					
Female Male					
5. Please indicate your ethnicity:					
Hispanic or Latino Not Hispanic or Latino					
6. Please indicate your race (mark one or more of the f	following):				
 American Indian or Alaska Native 					
Asian					
Black or African American					
Native Hawaiian or Other Pacific Islander White					
7. What is the highest degree or level of education you	ı have completed?				
High school graduate					
Completed some college					
Associate degree					
Bachelor's degree					
Completed some postgraduate Master's degree					
Ph.D.					
Medical degree					
Other advanced degree beyond a Master's degr	ree				
8. How long have you been involved with					
	1-5 years	6-10 years	11-15 years	16-20 years	20+ years
Early phase clinical trials	0	0	0	0	0
Cancer research	0	0	0	0	0

< Back

Your center

Save and Continue >

Your information will be saved every time you hit "save and continue". You may close and return to this survey anytime until it is submitted.

