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

National Cancer Institute

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

Introduction

The National Cancer Institute (NCI) and the [COOP GROUP NAME] would like your opinions about a new NCI clinical trial for [TRIAL DESCRIPTION].

They have developed a **brief online survey** tool to quickly and easily solicit feedback from physicians and research staff in the field to learn any potential issues with opening and accruing to this trial.

The PDF attachment to your email invitation provides an overview of the [TRIAL NAME] trial's concept. After reviewing this brief document we ask that you take 1 minute to answer this short survey.

Your comments will help us plan in advance for any concerns about the [TRIAL NAME] trial identified from this survey. We **thank you** for your assistance!

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

Next -->

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

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Privacy Statement and Consent

Your participation in this survey is completely voluntary. Please be assured that your participation in the survey is anonymous and your responses cannot be linked or associated with you.

You may skip any questions that you prefer not to answer. You are also free to stop participating at any point during the survey and have your responses deleted by clicking the "Opt out of survey" box at the top of each survey page.

This brief survey should only require approximately 1 minute of your time.

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

Next -->

Public reporting burden for this collection of information is estimated to average 1 minute 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 (0000-0000-000). Do not return the completed form to this address.

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

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Trial Summary Sheet

IMPORTANT:

Please review the 2-page trial description attached to the email you received regarding this survey.

You can open a copy of the document here.

(This document will open in a new tab.)

I have reviewed the trial description and am ready to begin -->

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

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	Opt out of survey
Please reply to all questions from the perspective	of <u>your</u> site.
Please indicate which best describes your site:	
My practice is located within an academic n	nedical center
 My practice is located within an NCI-Design 	nated Cancer Center
 My practice is located within a community h hospitals) 	ospital (i.e., non-academic, medical center
IWe are a free-standing private practice	
Other:	
1a. What best describes the size of your commi	unity hospital compared to others?
 We are a small-size community hospital 	(fewer than 100 beds)
We are a mid-size community hospital (b	
 We are a large-size community hospital 	
2. Please indicate which best describes your site's	s affiliation(s): (Please check all that apply)
CCOP	
□ MB-CCOP	
NCCCP	
ALLIANCE	
ECOG-ACRIN	
SWOG	
□ NRG	
□ COG	
□ EORTC	
COGNO (Cancer Australia)	
□ NCIC	
Other:	
^ 1M · L	
Which category best describes your role at your	r practice?
Physician	
Staff member/other	

Attachment_C10_prospectivesurvey

Note: Q1a appears only if "My practice is located within a community hospital" is selected for Q1.

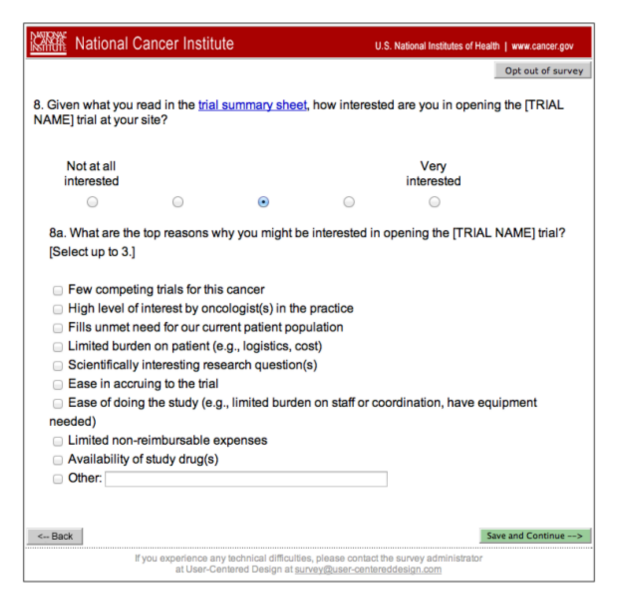
If user selects "physician" to Q3, continue below. If user selects "Staff member/other", continue HERE.

<u> </u>	ancer Institu	ıte	1	I.S. National Institutes of Health www.cano	er.g
				Opt out of	sur
What type of onco	logy best des	scribes your expe	ertise?		
Medical oncol	logy				
 Surgical onco 					
Radiation onc	7 2				
Gynecologic of					
Pediatric onco	1320 1300 1300				
[VARIABLE]	,,,,,,				
Other:					
0 001011					
entifically interest	ting it is to you]. [For each,] Please tell us hov	V
a. How scientific	ally interesting	ng is this research	n question to	you?	
Not at all				Very	
interesting				interesting	
0	0	0	0	0	
Please elaborate	<u> </u>				
RQ2: [INSERT R	Q2 FROM TE	RIAL SUMMARY	SHEETI		
			ACT OF THE PARTY.		
b. How scientific	ally interesting	ng is this research	h question to	you?	
	A)	(A)	1000	# T	
Not at all				Very	
interesting				interesting	
0	0	0	0	0	
0					
Plagea elaborata					
Please elaborate	:				
Please elaborate	:				
Please elaborate					
Please elaborate					
Please elaborate	:				

					Opt out of surve
. Assuming the [TF ndings will have or				hat impact do you	believe the
Low impact				High impact	
0	0	0	0	0	
The potential be to open and cor	nefits of this	trial for patient		e effort and resc	ources required
The potential be to open and cor Strongly	nefits of this	trial for patient		Strongly	ources required
The potential be to open and cor	nefits of this	trial for patient			ources required
The potential be to open and cor Strongly	nefits of this	trial for patient		Strongly	ources required

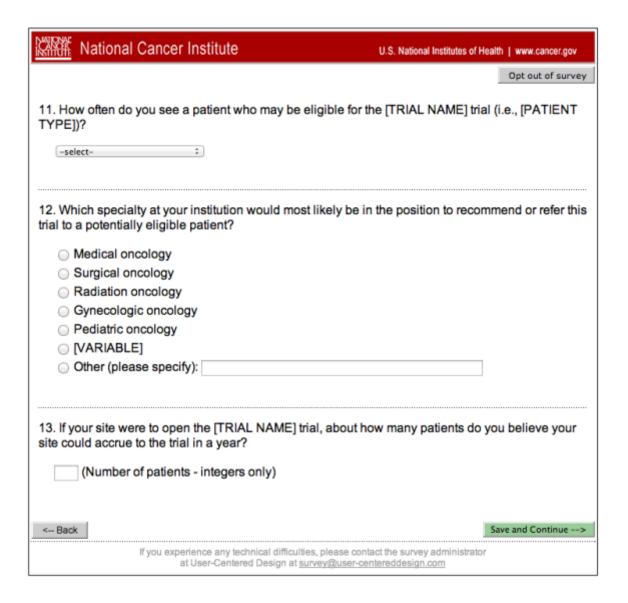
National (Cancer Institu	ıte	U.S	3. National Institutes of Hea	ith www.cancer.gov
8. Given what you NAME] trial at your		summary shee	et, how interested	d are you in openir	Opt out of survey
Not at all interested	•	0	0	Very interested	
8a. What are the [Select up to 3.]	e top reasons w	rhy you are not	that interested in	n opening the [TRI	AL NAME] trial?
☐ Too many co ☐ Limited inter ☐ Too difficult high refusals) ☐ Financial co		for this cancer st(s) here nts (e.g., rando	mization, screen	ning many to identit	
Does not maNot scientific	tch our patient ally interesting	population enough	den on staff, coo	ordination required,	equipment
needed) Other:		yaay (o.g., bar	3011 0111 012111, 0000	, amadon roquirou,	oquipmont
<== Back	,		ies, please contact th urvey@user-centered	e survey administrator	save and Continue>

Note: This 8a appears if the user selects either of the two leftmost radio buttons for $\mathsf{Q8}.$

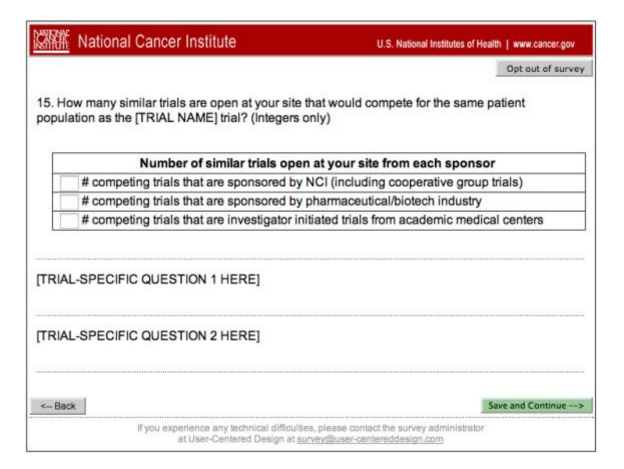


Note: This Q8a appears if the user selects any of the three rightmost radio buttons for Q9.

		ents listed below, p NAME] trial at you		l us whether		nake it
	ill this require un this trial at	ement make it to d your site?	open or	Not make it difficult to open/ run the trial	Make it somewhat difficult to open/ run the trial	Make it yery difficult to open/run the trial
[INSERT ITEM:	2 FROM TRIA	L SUMMARY SHE	ET]	0	0	0
[INSERT ITEM	N FROM TRIA	L SUMMARY SH	EET]	0	0	0
[INSERT ITEM	1 FROM TRIA	L SUMMARY SHE	EET]	0	0	0
[INSERT ITEM:	3 FROM TRIA	L SUMMARY SHE	ET]	0	0	0
[INSERT ITEM	4 FROM TRIA	L SUMMARY SHE	ET]	0	0	0
Overall, how dif	ficult do you b	elieve the [TRIAL I	NAME] tr	ial will be to		site?
0	0	0	0)	



How difficult will this issue make it to accrue patients to this trial?	Not make it difficult to accrue patients to the trial	Make it somewhat difficult to accrue patients to the trial	Make it very difficult to accrue patients to the trial
Burden on patient to participate in the trial (e.g., logistic time)	es,	0	0
Cost to the patient (e.g., insurance, reimbursement)	0	0	0
Patients declining to enroll (e.g., unwilling to randomiz prefer one study arm)	е, 💮	0	0
Explaining the trial's details to a patient, including consenting	0	0	0
Getting patients referred to the trial	0	0	0
Inclusion/exclusion criteria of the trial	0	0	0
	0	0	0





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Survey Complete	Opt out of surve
Thank you for comp	eting this survey!
Your answers have been s	ubmitted anonymously.
We appreciate your feedb Your comments will help ensure that we pla	
If you would like a summary of the findings afte 61164thflrlab@	
You may now clos	e this window.
If you experience any technical difficulties, i at User-Centered Design at <u>surve</u>	

END OF SURVEY

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	Opt out of survey
Please reply to all questions from the perspective of	your site.
1. Please indicate which best describes your site:	
 My practice is located within an academic me 	edical center
 My practice is located within an NCI-Designa 	ited Cancer Center
 My practice is located within a community hospitals) 	spital (i.e., non-academic, medical center
 I/We are a free-standing private practice 	
Other:	
1a. What best describes the size of your commun	6 5 5
 We are a small-size community hospital (fe 	
 We are a mid-size community hospital (be 	
 We are a large-size community hospital (n 	nore than 250 beds)
2. Please indicate which best describes your site's a CCOP MB-CCOP NCCCP ALLIANCE ECOG-ACRIN SWOG NRG COG EORTC COGNO (Cancer Australia) NCIC Other:	affiliation(s): (Please check all that apply)
Which category best describes your role at your p Physician Staff member/other	oractice? Save and Continue>
< Back	Save and Continue>

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Note: Q1a appears only if "My practice is located within a community hospital" is selected for Q1.

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	Opt out of survey
S1. What category best describes your role within your pract	tice? (Select one)
Research Nurse	
 Site Administrator / Manager 	
○ Coordinator	
○ CRA (non-nurse)	
Regulatory Specialist	
Data Manager	
Other:	
< Back	Save and Continue>
lf you experience any technical difficulties, please cont at User-Centered Design at <u>survey@user-ce</u>	

S2. Given what you read in at your site will be in opening. Not at all interested. S2a. What are the top read NAME] trial? [Select up to the property of the	asons why your sation 3.] resting enough patient population a patients (e.g., rasite would be too availability ct the study (e.g., trials for this can	site might not be andomization, scrugreat (e.g., non-burden on staff,	Very interested interested in oper	ning the [TRIAL dentify one; anticipatenses)
Not at all interested S2a. What are the top re NAME] trial? [Select up to Does not match our position of the Too difficult to accrue high refusals) Financial cost to our concerns about drug Too difficult to conduct needed) Too many competing Limited interest by on	asons why your sation 3.] resting enough patient population a patients (e.g., rasite would be too availability ct the study (e.g., trials for this can	site might not be andomization, scrugreat (e.g., non-burden on staff,	Very interested interested in oper	ning the [TRIAL dentify one; anticipatenses)
S2a. What are the top ren NAME] trial? [Select up to Not scientifically interest of Does not match our post of Too difficult to accrue high refusals) Financial cost to our selection of Concerns about druged Too difficult to conduct needed) Too many competing Limited interest by on	asons why your so as a site would be too availability ct the study (e.g., trials for this can	ndomization, scr great (e.g., non- burden on staff,	interested interested in oper	dentify one; anticipatenses)
S2a. What are the top reconstruction NAME] trial? [Select up to the NAME] trial? [Select up to the Not scientifically interested to the NAME] Too difficult to conduct the Name of Name	asons why your so as a site would be too availability ct the study (e.g., trials for this can	ndomization, scr great (e.g., non- burden on staff,	eening many to id reimbursable exp	dentify one; anticipatenses)
NAME] trial? [Select up to the content of the conte	resting enough patient population patients (e.g., rasite would be too availability of the study (e.g., trials for this can	ndomization, scr great (e.g., non- burden on staff,	eening many to id reimbursable exp	dentify one; anticipatenses)
Does not match our p Too difficult to accrue high refusals) Financial cost to our Concerns about drug Too difficult to conduct needed) Too many competing Limited interest by on	patient population patients (e.g., rasite would be too availability ct the study (e.g., trials for this can	great (e.g., non- burden on staff,	reimbursable exp	penses)
Other:	e or disagree with	h this statement?		
The potential benefits of to open and conduct it		atients are worth	Strongly	esources required
Disagree	0	0	Agree	
Please elaborate on your re	esponse above:			
< Back				Save and Continue

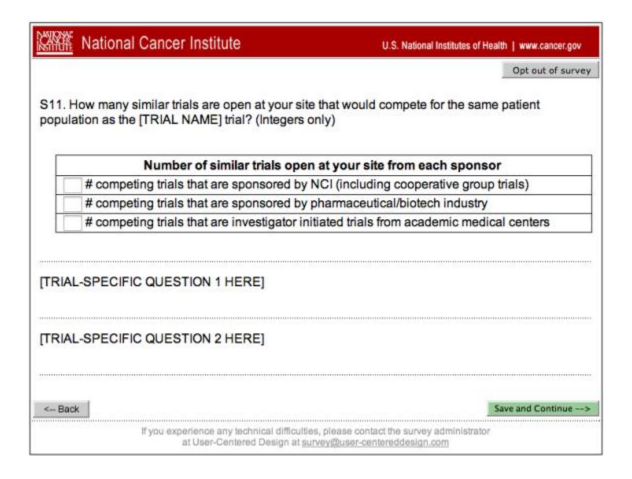
Attachment_C10_prospectivesurvey

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					opt out of survey
S2. Given what you at your site will be in				sted do you believe the	oncologists
Not at all interested				Very interested	
0	0	•	0	0	
S2a. What are the trial? [Select up to		hy your site m	ight be interes	sted in opening the [TR	IAL NAME]
Limited burde Fills unmet no Few competin Ease in accru Ease of doing needed) High level of Availability of Scientifically Other:	ing trials for this or uing to the trial g the study (e.g., interest by onco f study drug(s) interesting reserva- ou agree or disa	g., logistics, cont patient pop cancer limited burder logist(s) in the arch question(n on staff or co practice (s)	pordination, have equip	
Strongly Disagree				Strongly Agree	
0	0	0	0	0	
Please elaborate or	you experience any			Save	and Continue>

	to open/ run the trial	to open/ run the trial	open/run the trial
MMARY SHEET]	0	0	0
IMMARY SHEET]	0	0	0
MMARY SHEET]	0	0	
MMARY SHEET]	0		0
MMARY SHEET]	0	0	0
			site?
0 0			
al reasons – why you	believe this t	trial might be o	difficu
	MMARY SHEET] JMMARY SHEET] MMARY SHEET] MMARY SHEET] MMARY SHEET] e the [TRIAL NAME] to	MMARY SHEET] MMARY SHEET] MMARY SHEET] MMARY SHEET] MMARY SHEET] O The the [TRIAL NAME] trial will be to	MMARY SHEET] O O O O O O O O O O O O O O O O O O O

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		Opt out of survey			
S6. How often of [PATIENT TYP		ient who may be eligible for the [TRIAL NAME] trial (i.e.,			
-select-	:				
	cialty at your institution would tentially eligible patient?	d most likely be in the position to recommend or refer			
 Medical 	oncology				
 Surgical 	oncology				
 Radiation 	n oncology				
Gynecole	Gynecologic oncology				
Pediatric	 Pediatric oncology 				
○ [VARIAB	LE]				
Other (pl	ease specify):				
S8. If your site were to open the [TRIAL NAME] trial, about how many patients do you believe your site could accrue to the trial in a year? [Number of patients - integers only)					
< Back		Save and Continue>			
		difficulties, please contact the survey administrator gn at survey@user-centereddesign.com			

ur site? Not at all difficult			Very di	fficult			
\circ	0	0 0	0)			
S10. For each of the items below, please tell us whether or not you think it will make it difficult to accrue patients to the [TRIAL NAME] trial.							
How diff	icult will this issue patients to this		Not make it difficult to accrue patients to the trial	Make it somewhat difficult to accrue patients to the trial	Make it yery difficult to accrue patients to the trial		
Getting patier	nts referred to the tria	ıl	0		0		
	atient (e.g., insurance		0	0	0		
Explaining the consenting	e trial's details to a p	atient, including	0	0	0		
Inclusion/exc	lusion criteria of the	trial	0		0		
Burden on pa time)	tient to participate in	the trial (e.g., logistics,	0	0	0		
Patients decli prefer one stu		unwilling to randomize,	0	0	0		
Please elabor		itional reasons – why y	ou believe thi	s trial might b	oe difficult to		



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		Opt out of survey
S12.	Do you have any final comments about the [TRIAL	NAME] trial that you would like to share?
< Ba	ick	Submit Survey>
	If you experience any technical difficulties, please at User-Centered Design at <u>survey@use</u>	

