Date: Thursday, April 16, 2020 2:03:28 PM

View: SF - Reliance Information

Title: Reliance: Akron Children's Hospital -The Eating Disorders Quality Improvement Registry: A Feasibility Study

Reliance Information

* What Institution will rely on the Boston Children's IRB (the IRB ceding review to BCH)?

Akron Children's Hospital - FWA0000028600000917

If Other:

- 1.1 Please enter the name of institution.
- 1.2 FWA Number
- * Who is the Principal Investigator at the relying site?
 Maria Del-Pilar Trelles

If the person you need to add to your protocol cannot be found using the "Add" button above, please send an email to CHERP Support (cherp.support@childrens.harvard.edu) requesting that an account be created for the NON-BCH Principal Investigator. CHeRP Support will need the following information:

- First Name
- Last Name
- Email Address
- 3 What type of reliance agreement is being requested? Please select one:
 - 3.1 Smart IRB Master: Reliance agreement between BCH and another SMART IRB affiliated institution.

	See link for more information: https://smartirb.org/						
	3.1.1	What is the SMART IRB reliance application 1234	n ID?				
3.2		☐ Master (consortium-based): Reliance agreement among a consortium/network of institutions (other than SMART IRB).					
	3.2.1	Please specify consortium:					
3.3	.3 Other: Reliance agreement between BCH and another institution not affiliated with a master agreement						
	3.3.1	Please specify:					
Please only list researchers/staff engaged in the protocol at the relying site IF they have a conflict of interest(expect to have any financial interest, financial relationship, or position / advisory role with any other entity).BCH IRB considers 'engaged' to be interacting with subjects and/or obtaining individually identifiable data.							
Last	t Name	First Name	Employee ID	Role			
The	re are r	o items to display					
(cher	p.suppo	cannot be found using the "Add" button above ort@childrens.harvard.edu) requesting that an a CHeRP Support will need the following informations.	account be created for t	• •			
•	First I Last I Email						
expe entity	ct to ha y that n	Disclosure: Do any of the NON-BCH researd ave any financial interest, financial relations hay be affected by the research to be condu s providing funds or other support in conne	hip, or position or adv cted under the protoc	visory role with any other ol relationship with any			

5

O Yes No

If YE	S:					
5.1	5.1 Please describe the conflict of interest and any pertinent management plan.					
5.2	Please submit any pertinent documentation.					
	Name	Date Last Modified		Version	Owner	
	There are no	items to display				
Plea	se upload any	reliance request docum	nentation that the rel	ying site com _l	pleted, (if applicable)	
N	lame		Date Last Modified	Versio	n Owner	
F	RELIANCE EXA	MPLE.docx	4/16/2020 2:01 PM	0.01	Elizabeth Woods	
* Wil	I consent/asse	ent form(s) need to be ir	ndividualized for the	relying sites (include site specific	
		as additional HIPAA lanç n, addition of site heade	• •		, , , , , , , , , , , , , , , , , , ,	
	Yes No	ii, addition of Site neads	erriogo, etc. boller pr	ale signature s	section).	
* \A/:I	I	da a	- di: did. di - d f - u 41			
VVII	Yes () No	documents need to be in	ndividualized for the	relying sites?		
* Please indicate all research activities being conducted at the relying site and/or conducted by the relying site researchers at BCH. Check all that apply:						
~	Recruitment		,			
~	Consenting					
~	Medical Char	t/Record Review				
	Identifiable Da	ata Analysis				
~	Data Collecti	•				
	Other					
	J (1101					

Other:
Please specify:
Data Collection:
Please select one option:
Conducting surveys/questionnaires
O Drug/Device intervention
O Clinical exams and medical assessments (i.e. exams, x-rays, scans, EKG, ECHO, EEG, MRIs)
O Specimen collection (for clinical testing or research)
O Other
If Other: Please specify:

9.1 * Please describe all research activities being conducted at the relying site and/or conducted by the relying site researchers at BCH (specify where research activities will be conducted).

describe all research activities

10 What is your plan for communicating IRB actions, revised protocols, consents, etc. with this relying site (if different than described on the Multi-Site Information page, #1.2 of the main protocol)? plan for communicating

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Consents and Recruitment Materials

1 Please upload the site specific consent here. Please be sure that the title includes the name of the institution

Name	Date Last Modified	Version	Owner
RELIANCE EXAMPLE.docx	4/16/2020 2:03 PM	0.01	Elizabeth Woods

2 Please upload the recruitment documents here

Name	Date Last Modified	Version	Owner
RELIANCE EXAMPLE.docx	4/16/2020 2:03 PM	0.01	Elizabeth Woods

View: SF - PI's Statement

Title: Reliance: Akron Children's Hospital -The Eating Disorders Quality Improvement Registry: A Feasibility Study

PI's Statement

1 Upload any additional documents you think may be pertinent to this reliance request.

Name Date Last Modified Version Owner

There are no items to display

* I am aware of and support the reliance request that is being made for Boston Children's Hospital IRB to serve as the IRB for record for the above mentioned site. As the PI, I take full responsibility for submitting initial and ongoing information that requires IRB review from the other relying sites. I will also keep the relying site PIs informed about any associated IRB review activities and will make available to all sites the approved protocol, recruitment materials, consents, reports of actions, and any other documents and communications pertinent to IRB review. I assure that I have the appropriate resources to fulfill these additional responsibilities in order to assure all required human subject protection policies.

Yes No