

IRBManager User Guide for Researchers and Research Staff

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1.0 IRBManager

1.1 What is IRBManager?

IRBManager is an online submission, workflow, and data management system for the NMDP Institutional Review Board.

2.0 Login

2.1 Non-NMDP employees:

Existing user:

- 1. Go to https://nmdp.my.irbmanager.com/
- 2. Your User Name is your email address
- 3. Click on 'Forgot Password' if you need to reset your password
 - 3.1 You will receive an email with a link to reset your password

Note: If you do not use the email address associated with your account in IRBManager, you will not be able to reset the password. Should this occur, contact IRBStaff@nmdp.org in order to verify the email address on file. Do not create a new account. Duplicate accounts will not be merged. If you need to update your email address on file, contact <u>IRBStaff@nmdp.org</u>

New user:

Go to https://nmdp.my.irbmanager.com/

- 4. Click 'Click here to register'
- 5. Enter your email address on the first screen
- 6. You will be required to answer all the questions in the registration form
- 7. Once you register, you will receive an email with a link to reset your password. You must use this link within 24 hours. If you do not change your password within 24 hours, you will need to click on 'Forgot Password?' to have a new link sent.

2.2 NMDP employees:

- 1. If you are logging in to IRBManager for the first time, contact <u>IRBStaff@nmdp.org</u> to ensure you have access to the IRBManager Okta account.
- 2. Upon confirming access with IRB staff, go to https://nmdp.my.irbmanager.com/
- 3. Use your Okta login by clicking on the link next to Okta login

NMDP Users: To use your Okta login click here

For Help with Login please contact <u>IRBStaff@nmdp.org</u>.

2.3 How do I reset my password?

Click on 'Forgot Password?' to reset your password. You will receive an email with a link to reset your password. This link will expire within 24 hours. If you do not receive this email, check your spam/junk folder.

As a security precaution, IRBManager locks your account after three incorrect password attempts. It is recommended that you reset your password after two incorrect attempts. If your account is locked, use 'Forgot Password?' to reset your password and unlock your account.

3.0 Navigating the Home Page (also known as the DASHBOARD)

3.1 Actions

This is your Home/Dashboard screen. Use left-hand side Actions to start new xForms. For studies with which you are already associated, start a new xForm by first clicking on the study link under 'My Studies' (see section 3.6).

	Home						
PROGRAM*	My Studies						Take
Actions	Studies (1 Act	ive)					
Start a new Human Subjects Research Determination Form Start Initial Application for	• You are the	ciated with <u>1 active</u> Studies and <u>1 tota</u> Coordinator for <u>1 active</u> and <u>1 total</u> Stu	_				
Biomedical Study	xForms (2 Act	ive)					
Start Initial Application for Social and Behavioral Sciences Study	• You have 1)	insubmitted xForms. (Form being processed at a later stage. xForms awaiting your attention.					
Start xForm Show Sponsor Ids	• There are 1						
Recent Items							
2018-077-517	·	Only show events where I am: Vou have <u>4 Continuing Review</u> events.					
My Docs & xForms	You have <u>3</u> F	Request for Amendment events.					
0 Attachments 4 xForms	You have <mark>7 T</mark>	<u>otal Open</u> events					
	My Studies (1	Active)					
	Study	▼ Site	¢	PI	¢	Title	\$ Expires 4
	2018-077-517	Children's Hospital of Philadelphia		Tkachenko, Julia		July Test Study	08/29/2020

3.2 Messages

This heading is an area that the IRB will use for general communication to all the users within the system. Only displayed when there are current messages.

3.3 Studies

This area shows any studies with which the user is associated as Principal Investigator, Coordinator, Sub-Investigator, Administrative Contact, or other research staff. Click on the underlined link to view the details of the studies with which you are associated.

My Studies Studies (1 Active)

- You are associated with <u>1 active</u> Studies and <u>1 total</u> Studies.
- You are the Coordinator for <u>1 active</u> and <u>1 total</u> Studies.

3.4 xForms

IRBManager uses web-based forms called xForms. The xForms section shows the forms currently being processed at this time for the user.

xForms (5 Active)

- You have <u>2 unsubmitted</u> xForms.
- You have <u>3 xForms</u> being processed at a later stage.
- There are <u>1 xForms</u> awaiting your attention.

In the above view of the DASHBOARD:

- You have 2 unsubmitted xForms.
 - The user currently started 2 xForms (submissions) but clicked "Save for Later" or closed out of the form without submitting the form.
 - Note: By clicking the link, '2 unsubmitted', the user can go straight to the unsubmitted xForms.
- You have 3 xForms being processed at a later stage.
 - The user has submitted 3 forms that are currently awaiting review and signature by the PI or are being processed by the IRB.
 - Click on the '3 xForms' link to see the current stage of each xForm
- You have xForms in error (not shown in screenshot).
 - The user has some forms that have errors. This is a software issue, and the user must notify the IRB office if they receive this notification.
 - $\circ~$ Note: If the user has an xForm in error, the user can click the underlined link and go straight to the xForm in error.
- There are 1 xForms awaiting your attention.
 - The user has 1 xForms that currently need action by the user (e.g., review and signature required, revisions requested by the IRB).
 - Note: By clicking the link, '1 xForms' the user can go straight to the xForms that need the user's attention.

3.5 Events

The events section of the DASHBOARD shows the total number of open events and the number of events categorized by name, such as Continuing Review, Initial Submission, or Amendment. The drop-down following 'Only show events where I am:' will allow you to show the events according to your role in the associated study.

Events (11 Open)	
 Only show events where I am: You have <u>8 Continuing Review</u> events. You have <u>3 Request for Amendment</u> events. You have <u>11 Total Open</u> events 	

3.5.1 Event Detail

vent Details: Request	for Amendment on 2018-07	77-517			Help Submitter		
Study-Site							
Study:	2018-077-517			Site: 517 - Childre	n's Hospital of Philadel		
Title:	July Test Study		Comn	nittee: IRB	a: IRB		
PI:	Tkachenko, Julia		Spon	sor Id			
Event							
Туре:	Request for Amendment		St	arted: 09/16/2019			
Instance:	No		Comp	leted:			
Committee:	Inherited from Study						
Review Type:	Expedited						
Action Date:	9/16/2019		Additional IR Comm	B NoA nents:			
Steps (9)							
Step		Planned	Actual	Complete	Minutes		
Receive Request for Am	endment		09/16/2019	Yes			
Staff Review			09/16/2019	Yes			
Administrative Review			09/16/2019	Yes			
Place Expedited Approva	al on FBA	09/30/2019	12/28/2019	Yes	12/28/2019		
Notify PI of Outcome				No			

By clicking into the event, you can view event details:

- Study: Study IRB number
- Site: NMDP Network Number (if available) and the name of the institution
- **PI:** Principal investigator's name
- **Type:** the type of event, which could include Initial Submission, Continuing Review, Adverse Event/Unanticipated, etc.
- **Started:** the date the event first started
- **Steps:** each event will have the steps listed. When the NMDP IRB has completed these actions, the complete column will be marked 'yes'.

3.5.2 Event Actions

Actions in the event details screen on the left-hand side are as follows:

- Attachments: shows the attachments related to the event detail
 - $\circ~$ Generated Docs shows the approval documents associated with the study
- **xForms:** shows the form used to create the event.

• **Done:** returns the user to the next most logical page depending on previous actions.

3.6 My Studies

My Studies (1 A	ctive)						
Study \$	Site +	PI \$	Title \$	Expires \$	Status -	Reference Doc(s)	÷
2018-077-517	Children's Hospital of Philadelphia	Tkachenko, Julia	July Test Study	08/29/2020	Open to accrual		

This is a list of all the active studies and site(s) with which the user is associated. Note: A study is not created under My Studies when a user submits an xForm. A study is created when the IRB determines that the study is ready for review. From your DASHBOARD, you can view the following information regarding your studies:

- The study number.
- The site where the project is taking place.
- The PI name.
- The study title.
- Expiration date of the project (if blank study has not yet been approved).
- Status: This will depend on whether the study is still under initial review or if elsewhere within the project's lifecycle. Statuses can include Active, Closed to enrollment, Data analysis, Data collection, Disapproved, On hold, Open to accrual, Pending Approval, Study Completed, Terminated, or Withdrawn.
- Reference Doc(s): will display documents that have been approved by the IRB. Note: this field will likely be blank as Reference Doc(s) are not being populated on the study-sites. Approved study-site documents will be displayed under Attachments (see section 3.6.2.1).

By clicking the small arrow to the right of Project, Site, PI, Title, Expires, and Status, you can sort the studies accordingly.

- Study can be sorted by Project Number
- Site can be sorted alphabetically
- PI is sorted alphabetically
- Title is sorted alphabetically
- Expiration dates are sorted by numerical date
- Status is grouped and sorted alphabetically

3.6.1 Study-Sites

Click on the blue link under "Study" (see 'My Studies') to view study-site information and make submissions associated with that study-site.

Home :udy 54321Test-916 (IF	(B)		
Study:	54321Test	Sponsor(s):	NIH (Primary)
Committee:	IRB	Sponsor Id:	1234
Category:		Grants:	
Department:			
Agent Types:		CRO:	
Title:	Test Study	Year:	2020
FDA Expedited Categories:		FDA Regulated:	No
	Adults with impaired decision-making capacity are allowed to be enrolled on this protocol.	Minor Participants:	N/A
NMDP IRB Review Responsibility:		Number of Participants Approved:	
Protocol ID:	123	Protocol Version:	2.0
Research Category:	Blood for Lab Research		
tudy-Site			
Site(s):	916 - 916Test	PI:	Tkachenko, Jul
Status:	Active	Additional:	N
Approval:	June 6, 2022	Expiration:	N/A
Initial Approval:	June 6, 2022	Other Expirations:	
Is Primary Site?:			
Comments:	Example site		

The main/study-level IRB approval information is listed under a MASTER site. All other study-site information is listed under study-site section. This section notes information about the study and the study-site, such as:

- Site(s): NMDP Network Number (if available) and the name of the institution
- PI: Principal Investigator of the study at the site
- Status
- Approval
- Initial Approval
- Expiration
- Comments

3.6.2.1 Attachments

Beginning June 27, 2023, final documents approved by the NMDP IRB for study-site submissions will be added as attachments to the study-site under study-site attachments.

 Study-Site A 	 Study-Site Attachments (3) 				
Action	Name +	Туре +	Date 🔸	By \$	
🛃 🖼 🗒 🔝 🗶	Test Consent Form 1	Consent Form	06/27/2023	jtkachen@nmdp.org	
🛃 🖼 🗒 🕅 🖉 🗙	Test Consent Form 2	Consent Form	06/27/2023	jtkachen@nmdp.org	
🚰 👒 🔚 🐚 🔣 🗙	Test Consent Form 3	Consent Form	06/27/2023	jtkachen@nmdp.org	

3.6.2 Study-Site Contacts

Study-site contacts are directly associated with a specific site on the study. They have access to make submissions specific to their study-site. This section lists the contacts associated with the study.

Study-Site Contacts (2)				
Name •	Role \$			
Jobe, Christina	Principal Investigator			
Test, Submitter	Coordinator			

Current study-site contacts can add or remove other study-site contacts:

- Select a study under 'My Studies' on the homepage.
- Click on 'Start xForm' on the left hand-side under the Actions and select the Add/Remove Study-Site Contacts xForm. If the new contact's email is not populating, the xForm will have an option to click on a link to add a new contact to the system.

3.6.3 Study Contacts

Study Contacts are comprised of the central study-level protocol team and can be listed under the main study details (separate from being listed as a study-site contact under the MASTER study-site). Study Contacts have access to the IRB-approved study sites and can view existing Events and submissions for approved participating sites. They can also see pending xForms, what stage they are in, and who started them for already approved study-sites.

Study Contacts can see all the study-sites they are associated with under 'Studies':

Studies (16 Active)

- You are associated with **<u>16 active</u>** Studies and **<u>17 total</u>** Studies.
- You are the Primary Contact for <u>**1** active</u> and <u>**1** total</u> Studies.
- You are the Research Staff for <u>1 active</u> and <u>1 total</u> Studies.
- You are the Study Contact for <u>15 active</u> and <u>16 total</u> Studies.

*active means approved by the NMDP IRB

For study-sites that are **approved** (active) by the NMDP IRB:

• To view study-site activities, select the site from the list of 'My Studies'. Pending xForms will appear under the left-hand side Actions menu or under the corresponding Event. Note: an Event is created when IRB Pre-Review is completed and the xForm has moved to the next stage (see section 5.1)

For study-sites that are **pending** NMDP IRB approval:

- The study-site is created when the IRB Pre-Review is completed and will appear under 'Studies'. Note: 'My Studies' only shows sites that have already been approved.
- To view the site's Initial Submission prior to the IRB Pre-Review, the study contact will need to ask the submitter to add them as a collaborator on the **NMDP sIRB Study-Specific Local Context Worksheet** xForm (see section 4.1.2).
- After the IRB Pre-Review is completed, the xForm can be viewed under the Site Initial Submission Event.

To add or remove a study contact, contact IRBStaff@nmdp.org

3.7 Find Study

Users can search by PI name or complete NMDP IRB study number (e.g., IRB-1991-0002) to find a study with which the user is associated. Note: A study is not created when a user submits an xForm. A study is created when the IRB determines that the study is ready for review.



3.8 Help

Under **Help** you can contact the NMDP IRB Staff by choosing 'Contact Administrator'. Page help is currently disabled. Use this manual for assistance with using IRBManager.

3.9 User's Settings

Allows the user to edit their profile settings. By clicking on your Settings, you will access My Settings.

Under My Settings you can:

- 1. Change your password
- 2. Change your profile
- 3. Change your phone number
- 4. Change your address
- 5. View last 25 logins
- 6. Create an email signature

3.9.3 Change My Password

1. Step 1: Click on "Change My Password" link. You will then see this view:

Change Password	
User	Test Generic Coordinator Login
Username	Coordinator
Old Password	
New Password	
New Password Confirm	
	Update

- 2. Enter old password.
- 3. Enter new password that meets requirements and then enter new password again to confirm.

Your password must meet the following requirements:

- 1. Be 6 or more characters
- 2. Be different from the username
- 3. Contain characters from at least 3 of the following: upper case letter(s), lower case letter(s), number(s), and special characters (e.g. !, @, #)

4. Click "Update".

3.9.4 Change My Profile

1. Click on "Change My Profile" link. You will then see this view:

My Profile	
Change User Informat	tion
First Name	Test Generic
Last Name	Coordinator Login
Degree	
Specialty	
Email Address	
	Update

- 2. Update your name, degree, specialty or email address.
- 3. Click "Update."

3.9.5 My Phone Number(s)

1. Click on "My Phone Number(s)" link. You will then see this view:

	DI	Margare Income
мγ	Phone	Numbers

Numbers				
Action	Primary	Туре	Number	
🚰 🗙	✓	Business	3-2693	

2. Click the icon that appears to be a hand holding a piece of paper.

You will then see this view:

Contact Name	
Full Name:	Test Generic Coordinator Login
Phone Book	
Phone Type:	V
Primary Number:	
	Add Phone Cancel

- 3. Update or add your phone number.
- 4. Click "Update Phone." Or "Add Phone".

3.9.6 My Address(es)

1. Click on "My Address(es)" link.

You will then see this view:

Address Book (1)								
Action	Primary	Туре 🔹	Street \$	City \$	State \$	Zip 🗢	Country	
ピ 🗙	\checkmark	Business	10 Anywhere St	Anywhere	California	11111	United States	

2. Click the "hand holding a piece of paper."

You will then see this view:

Contact Name			
Full Name:	Investigator, Test MSc	Email:	test@oshpd.ca.gov
Address Book			
Address Type:	Business 🗸	Country:	United States
Street:	10 Anywhere St		
Line 2:			
Line 3:			
City:	Anywhere		
State/Province:	CA - California	Zip/Postal Code:	11111
Primary Address:			
	Update Address Reset Cancel		

- 3. Update your address.
- 4. Click "Update Address."

3.9.7 Last 25 Logins

Clicking on this link will provide a list of the time and date of the last 25 times you logged into IRBManager.

4.0 Managing forms in IRBManager (xForms)

Users submitting to the NMDP IRB will submit the xForm appropriate to the type of request (e.g., new study, amendment, continuing review). xForms are started under the action 'Start xForm' from either your Dashboard or a specific study or site. Read the description next to each xForm to determine which one to use.

1. The xForms below are started from your Dashboard. On your home/dashboard, click 'Start xForm' on the left and then choose the appropriate xForm.

- Application for Multi-Site Coordinating Center Activities
 - To be completed along with the Initial Application for federally funded multisite studies if the Coordinating Center is not already approved by the NMDP IRB.
- Human Subjects Research Determination Form
 - This form should be completed when the investigator is uncertain whether the study is human subjects research needing NMDP IRB review.
- Initial Application for Biomedical Studies
 - This application should be used for new bio-medical studies.
- Initial Application for Social and Behavioral Sciences
 - This application should be used for new social and behavioral sciences studies.
- NMDP sIRB Study-Specific Local Context Worksheet
 - This application should be used for study sites needing to open a study using NMDP single IRB.
- Request to Rely on an External IRB
 - Use this form to request that an NMDP-initiated study relies on an external IRB. NMDP will only rely on an external IRB that is accredited by the Association for the Accreditation of Human Research Protection Programs (AAHRPP).
- Single IRB Signatory Institution Enrollment and Local Context Form
 - Use this form when your institution is enrolling in the NMDP single IRB to provide contact and local context information for your institution. This form is completed once by the institution's IRB/HRPP staff.

2. The study-specific xForms below are started from your study-site. First click the link for the study under 'My Studies'; then click 'Start xForm' on the left and then choose the appropriate xForm.

- Add/Remove Study-Site Contacts
 - Use this form to add or remove any study-site contacts, other than the Principal Investigator.
- IRB Request for Study Amendment
 - Relying institutions should use this form to submit revised consent/assent forms, translations, and other participating site study specific materials (not short forms). Main study level teams (not site specific) should use this form for changes to the protocol and other study-level materials, as well as translations and overall/main PI changes.
- Major Protocol Exception Request form

- A one-time, temporary departure from the IRB-approved protocol procedures that is identified before it occurs, intended for one specific study subject, and may adversely affect the safety, rights, or welfare of the subject(s) and/or the scientific validity of the research.
- Notification of Study Closure
 - Notification of Study Closure form for the Study-Wide closure at the study level (not site specific).
- Reportable Event Form
 - For reporting potential unanticipated problems involving risks to subjects or others and potential serious or continuing non-compliance to the NMDP IRB for review.
- Revised NMDP sIRB Study-Specific Local Context Worksheet
 - Relying institutions should use this form to make any changes to the previously submitted information on the NMDP sIRB Study-Specific Local Context Worksheet including study-site PI.
- Study Site Closure Form
 - Relying institutions should use this form to close a study at their site prior to the study closing as a whole.
- Temporary Variance Request Form
 - For study level (not site specific), an intentional change to the previously IRB-approved protocol that is initiated by the sponsor/Study Chair and is implemented as a systematic change prior to an official protocol amendment.

3. The NMDP sIRB Site Management xForms below are started from your Site Management page (see section 6.0).

- Add/Remove sIRB Site Contacts
 - $_{\odot}$ $\,$ Use this form to add or remove any contacts on your sIRB Site Management.
- Annual Renewal or Updates to sIRB Enrollment and Local Context Instructions
 - Instructions for completing annual renewal or updates to the sIRB Enrollment and Local Context Form.
- Site-specific Documents Submission
 - Use this form to submit your site-specific documents that are not specific to a study (e.g., Short Form templates, and other research participant-facing template documents that can be used for any study.)

4.1 Completing an xForm

The questions in the xForms are directed by the answers given. For example, if you mark that the study does not involve NMDP donors, the form will not ask subsequent questions about NMDP donors. Changing responses on the form will enable subsequent questions that were hidden or hide questions that are no longer applicable.

To navigate an xForm, click "Next" to move on to the next page of the xForm. Click "Save for Later" to stop working on the form. The xForm can then be reopened by going to the xForm link under "Actions."

Previous Next Save for Later More	►
-----------------------------------	---

Most questions on the form are required (the question will be labelled "Required"). All required questions on the form must be completed before submitting. If they are not, a red error message will appear. The system will not allow you to continue until the field has been populated.

All required questions on a page must be completed before clicking 'Next'. To skip to another page on the form without answering all questions on the page at that time, use the navigation drop-down at the top of the page.

MATCH	a Collaborators	Study Information	Page 1 of 20			
n for Biomedical Studies Study Information						
NMDP IRB Study Number	N/A					

The jurisdiction of the NMDP IRB for this study is solely for the protection of the rights and welfare of unrelated dor Be The Match Registry as research subjects. The protection of any other subject population on this study (e.g. trans related donors) is the jurisdiction of the transplant center's IRB.

4.1.1 Adding Contacts

Some questions ask for a study contact. Type in the name or email address of the contact and choose the appropriate contact from the list that appears. If the contact does not appear, they are not included in the system. Click on the link provided to add a contact to the system:

Please enter the email address of the Principal Investigator.	(Required)	Add Note	View Audit
If the Principal Investigator has not been entered as a conta Click here to add a new contact to the system	ict, please use the link below to enter a new contact.		

A new tab will open. Enter the required information for the new contact. Click 'Submit' to activate the new contact. Note: this action only pertains to the Contact Form. You may click 'Submit' or 'Save for Later' to continue working on the main application. In a moment, you should be able to select the new contact from the drop-down list.

4.1.2 Adding Collaborators

Collaborators can be added to an xForm to view, edit, manage, or submit the form. To add a collaborator, click on the 'Collaborators' link on the top of the page:

	Collaborators	Protocol Identification 	Page 1 of 6	Nex
NMDP sIRB Study-Sp	ecific Local Context Worksheet	- Protocol Identification		
Submitte	er		Add Note	View Audit
Test, Sub	omitter			
	Email: abc@yahoo.com	Phone:		

4.1.3 Attaching a Document

To attach a document:

1. Click "Add Attachment" on specified attachment questions.

- 2. Do not change the name of the document. The NMDP IRB Notice of Action will display the file name of the documents as the name of the attachements.
- 3. Select the type of attachment if that option is available. On some questions, only a certain type of document is intended to be attached, and the type is preset.
- 4. Click the "Select" button to locate the file on your computer (files can also be attached from Dropbox or Box)
- 5. Click "Attach" to finish the process.

Add Attachme	ent
Name:	
	(leave blank to use name of uploaded file)
Туре:	Assent
File:	Select
	Attach Cancel

Note: additional documents can be attached to any question. Click 'add note', save, then click on attachment icon to add attachments.

4.2 New study applications

From your Dashboard, under the action menu on the far-left side, select one of the following:

- 1. Start an Initial Application for Biomedical Study **or** Start Initial Application for Social and Behavioral Sciences Study (These applications should be used for new studies.)
- 2. Start xForm
 - a. Gives the option to start one of the above forms or to start the 'Add New Contact to IRBManager' form, which is used to create a username and login for a contact not in the IRBManager system.
 - b. Click on the name of the form to start the form.

4.2.1 Submitting an xForm

Before submitting an xForm, review the form completely. Changes cannot be made once an xForm is submitted, unless the IRB or PI requests changes.

After all pages on the form have been completed, the submission screen will appear. "Submit" will need to be clicked to complete the submission.

Form Completed						
You've completed the form. You can now either save the form for later revision, or submit it.						
Go Back Save for Later F	Print Submit					

If you are not the PI, the form will go to the PI for signature. They will receive an email from IRBManager indicating their review and signature is required; the email will contain a link that will take them to the form. If the PI wants to return the form to the submitter, they will have the opportunity; otherwise, the PI signs and submits the form.

Note: If the PI does not see the email from IRBManager, they should check their Spam/Junk folder and make sure to mark emails from IRBManager as safe.

The PI can also access the form from their Dashboard. Under the xForms section of their Dashboard, they will see an xForm awaiting attention:

x	xForms (2 Active)				
0	You have <u>1 unsubmitted</u> xForms.				
0	You have <u>1 xForm</u> being processed at a later stage.				
0	There are <u>1 xForms</u> awaiting your attention.				

By clicking on the hyperlink to the xForms, the study PI is able to see the forms, click on the study hyperlink, and review/sign the form.

Form	Identifier [‡]	Owner \$	\$\$	Status 🕈	\$	Requires Approval ^{\$}
Initial Application for Biomedical Studies	Christmas Study	Tkachenko, Julia (jtkachen@nmdp.org)	PI Signature for Research Staff Submission	Work in progress	10 minutes ago	No

4.3 Requesting PI Signature (not required for some xForms)

To request a change, the PI can click on 'Add Note' next to the question, enter comments, and save:

their spouses	dy Chair or any Principal Investigator involved in the development or coordination of this study (or Add No s or dependent children) have a Significant Financial Interest related to his/her Institutional Responsibilities t s/her objectivity?	
	Enter Note:	ا
	Change to Yes	
	Save Cancel	
No		

The PI will select either 'Ready for Submission' or 'Revisions required prior to Submission.' If Revisions are required, the PI will be prompted to indicate revisions:

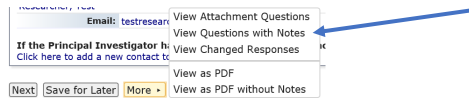
Is this Initial Application ready for submission?	(Required)	
Ready for Submission		
Revisions required prior to Submission		
Please indicate any revisions required prior to a	submission. (Required)	480
		ABC

They will click Next and then Submit

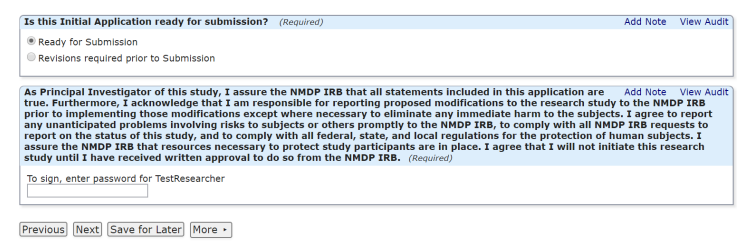
4.3.1 Making requested revisions to the xForm

If the PI has indicated that revisions are required, the form will go back into the Data Entry stage. The submitter will receive an email with revisions required. Click on the link in the email or access the xForm from your dashboard.

To view PI notes, click on 'More' button and select 'View Questions with Notes'



After making changes, click Submit. The form will go back to PI for signature. If the form is Ready for Submission, the PI will sign the form by entering their IRBManager password:



The form will now be submitted to the NMDP IRB office for review and will move into the 'IRB Pre-Review' stage.

4.3.2 Clarifications required by the IRB Staff

If IRB Staff requires clarifications, you will receive an email with Clarifications Required. Click on the link in the email or access the xForm from your dashboard to respond to the clarifications.

If there are any specific questions requiring clarifications, click on 'More' button at the bottom of the form and select 'View Questions with Notes.' Otherwise, navigate to 'Respond to IRB Questions' to provide a response:

a Collaborators	Study Information	*	Page 1 of 23
Study Information	Study Information		
A.	Institution Information		
	Funding		

4.4 Study Submissions

Study specific forms such as Request for Amendment and Notification of Study Closure, must be started from within the study. Navigate to the study. On the left-hand side under Actions, select Start xForm:

	Home	
PROGRAM*	Study 2018-077-517 (1	(RB)
Actions	▼Study	
Send EMail	Study:	2018-077
Start xForm	Committee:	IRB
xForms (0)	Category:	
Recent Items	Department:	
2018-077-517	Agent Types:	
IRB-2020-0016-506	Title:	July Test Study
	Coordinating Centers:	
Useful Links	FDA Expedited	

Click on the Form you wish to submit and begin completing the form.

After all pages on the form have been completed, the submission screen will appear. "Submit" will need to be clicked to complete the submission. If you are not the PI, the form will go to the PI for signature. They will receive an email indicating their review and signature is required. If PI wants to return the form to the submitter, they will have the opportunity; otherwise, the PI signs and submits the form.

If IRB Staff requires clarifications, you will receive an email with Clarifications Required. Follow instructions above.

5.0 IRB Determinations

The IRB has completed the review, and the IRB Office is preparing the decision notification to the research team.

5.1 Checking the status of an xForm

Click on xForms under your Home screen or 'xForms' under 'My Documents & Forms' on the left-hand side of the screen to see a list of all xForms with which you are associated (either as a PI, administrative contact, coordinator, etc.).

Home				🙊 Find Study (Ct	rl+Q)	2
4y Forms				Help Submi	tter's Settings S	Sign off
				Filter:		
i	What's this?					
This	s table shows xForms associated with you as	a user of IRBM	lanager across th	e entire system.		
	Status: All	•				
Action	Form +	Identifier +	Owner A			
			Owner -	Stage \$	As Of	
×	Human Subjects Research Determination Form	sdf				• 24 PM
×			Test, Submitter (testsubmitter)	Human Subjects Research Determination Data Entry		

1 completed xForms are Hidden.

- Action: If a form can be deleted or copied for an amendment, those options will display in this column. Note that an xForm can only be deleted when in the Data Entry stage.
- Form: Name of the xForm
- Identifier: This is generally the PI associated with the project
- **Owner:** The owner of the form will be the user who started the xForm until a project is created (a project will be created within IRBManager after the IRB has sent the project for review), and then the form is owned by the project
- **Stage:** This is the current stage that the form is in. Stages will vary by form, but in general will be the following:
 - $\circ~$ Data Entry: Form is currently with the user/submitter either to complete or make changes.
 - PI Signature for Research Staff Submission: If the PI is not the submitter, this is the stage where the PI reviews and submits the form.

- IRB Pre-Review: Stage that the IRB Office will screen the application for completeness and either request further information/documentation (clarifications) or assign for review. Note: this stage may include Administrative review and approval, if applicable.
- Expedited Review: The project is being reviewed at the expedited level.
- $\circ~$ Full Board Review: The project is being reviewed by the convened board.
- Final Processing: The IRB has completed the review, and the IRB Office is preparing the decision notification to the research team.

You can also check the status of the xForm from within the event (see 3.2 for finding an xForm on an event). Note: the xForm will be 'owned' by the type of event with which it is associated (i.e. initial submission, continuing review, amendment [once the amendment is submitted]).

5.2 IRB Decision Notification

Once the IRB Office or the IRB reviewer(s) have completed their review, the research team will receive an email detailing the IRB decision. The appropriate approval/determinaton letter will be attached.

- **Approved**: No changes are requested by the IRB; the investigator may initiate the study or continue the study, or may implement the changes to the protocol.
- **Approved with Stipulations**: Study may <u>not</u> be initiated until Stipulations are met.
 - The form will go to the Data Entry stage. If changes are requested to specific documents, please replace current documents with red-lined copies. Click on the green arrows to replace a document, rather than deleting the document and adding the new one. State your response in the following question of the xForm 'Please use this box to respond to any IRB questions, if needed'. Alternatively you may attach a formal response letter with any additional attachments under 'Add additional attachments, if needed'. Once the revisions are submitted, the form will either go back to the primary and secondary reviewers or back to the Full Board. If additional changes are requested, the form will return to Data Entry for revisions are met.
- **Disapproved:** The research is not approved.
- **Deferred:** Substantive clarifications or modifications are required by the IRB before the research may be initiated or continued beyond the current approval period, or before the protocol change may be implemented. The convened IRB must review the response to the deferral.
 - Form will go back to data entry and be routed back through the IRB Office and to the full board for review.
- **Tabled:** If the convened IRB is unable to adequately review a study due to lack of time or loss of quorum, the study will be reviewed at the next full board meeting.

6.0 Using the NMDP IRB as a Single IRB

Signatory Institutions relying on the NMDP IRB have a Site Management page in IRBManager. This page is created when a site enrolls in the NMDP single IRB (sIRB). Primary contacts associated with a Signatory Institution have access to their site under `My Studies':

My Studies								Help	Submitter's Settings Sign o
My Studies (97 Active)								
Study 🔸	Site	\$ PI	¢	Title	¢	Expires	¢	Status +	Reference Doc(s)
Site Management- Test Site	Test	Tkachenko, Julia		sIRB Site Management		Annual Renewal Required		Enrolled	Boilerplate ICF ACH Template

Click on the blue link under "Study" to view site information and make submissions associated with the site.

Home											Find Stu	ıdy (Ct	rl+Q))		8
NMDP sI	RB Site Manageme	nt-Te	st Site (Single IRB Institutions)								Help	Subn	nitte	r's Setti	ngs S	ign o
	sIRB														\$	collaps
NMDP sIRB: Site Management							Title:	sIRB	Site Manag	em	ent					
NMDP s	IRB-Site															
	Site(s): Tes	st Site	e - Test		Si	gnato	ry Official:	Heir	nen, Mirano	da						
	Location(s):															
Status: Enrolled							Additional:	N								
Approval: September 14, 2021							Expiration:	Ann	ual Renew	al F	Require	d				
Initial Approval: September 14, 2021					Ot	her E	xpirations:	sIRE	8 Enrollment	: Re	newal D	ate - (09/20	/2021		
Affili	ate Institutions: Affi	liate 1	. FWA#1234567		Compone	ent Ir	stitutions:	Com	ponent 1							
	Comments: Sep	otembe	er 2021													
 Site C 	Contacts (3)														9	collaps
Name				•	Role											٠
Jobe, Ch	ristina				Primary Contact											
Test, Sul	omitter				Primary Contact											
Tkachen	ko, Julia				Primary	Cont	act									
	sIRB-Site Referer	nce Do	ocuments (1) 🗏												9	collaps
Name					\$	Тур	e				• Acti	ve		• Inac	tivate	d 4
10-CBA	Recipient Consent v5	.0_rev	091321_FINAL 09-13-2021.docx			Boile	erplate Conse	ent L	anguage.		09/2	1/202	1			
 Refer 	ence xForms (1)														\$	collaps
Action	Form					•	Stage	• A	s Of	¢	Ref Ac	tive	¢	Inactiv	ated	
	Single IRB Signator	ry Inst	itution Enrollment and Local Context Form				Complete	0	9/21/2021		09/21/	2021				
 Event 	s (3)														\$	collaps
Event	¢	Att	Instance/UDF									÷	Sta	rt 🔺	Comp	olete +
Study-Si	te Personnel Change	0	Replace Signatory Official Add new Primary	Contact									06/	06/2023	06/06	/2023
sIRB Rer	newal	1											09/	21/2021		

Contacts listed under Site Contacts act as the point of contact for their institution. They can submit annual sIRB review, update their boilerplate template language, and request other site documents to be approved (see section 4.0).

For further information about sIRB, reference **NMDP Single IRB Manual for Local Institutions** found on the NMDP IRB website

https://network.bethematchclinical.org/research/institutional-review-board/relying-on-the-nmdp-sirb/

Further Questions

If you have further questions regarding your study status or using IRBManager, contact the NMDP IRB Staff at IRBStaff@nmdp.org.