TABLE OF CONTENTS
*Page 2-5: The Institutional Profile

*Pages 6-11: Local Considerations: Study-specific Human Research Protections Reviews (completed by the HRPP Liaison)

*Pages 12-16: The PI Survey regarding Conduct of the Study at their Site
The Institutional Profile
(sections 1 and 2 pertain to local considerations)
ABOUT THE INSTITUTIONAL PROFILE
The IREx IP contains basic information about your institution that another institution might use to determine if they feel comfortable relying on, or serving as the Reviewing IRB for, your institution. The IP can also be used by IRBs and study teams to better understand one's institutional processes and requirements when using reliance.

This information will be visible to other users in IREx and publicly available as a downloadable PDF on the IREx Website here. This information is for general review purposes only. It may not be accurate and may be subject to change, withdrawn or revised at any time without notice. The Institutional Profile is not intended to serve as a complete record of an institution’s study-specific local considerations.

Section 1: GENERAL HRPP INFORMATION

<table>
<thead>
<tr>
<th>Institution</th>
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<table>
<thead>
<tr>
<th>Federalwide Assurance (FWA) #</th>
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</thead>
<tbody>
<tr>
<td>(Enter FWA# and 8 digits, eg FWA#00000000)</td>
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</table>

<table>
<thead>
<tr>
<th>FWA Expiration Date</th>
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<tr>
<td>___________________</td>
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</table>

<table>
<thead>
<tr>
<th>Does your institution have an internal IRB?</th>
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<tbody>
<tr>
<td>☐ Yes</td>
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<tr>
<td>☐ No</td>
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<table>
<thead>
<tr>
<th>IRB Registry Number(s)</th>
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<tr>
<td>_______________________</td>
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<table>
<thead>
<tr>
<th>Is the IRB AAHRPP accredited?</th>
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<tbody>
<tr>
<td>☐ Yes</td>
</tr>
<tr>
<td>☐ No</td>
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</table>

<table>
<thead>
<tr>
<th>Does your institution REQUIRE institutions to be AAHRPP accredited in order to serve as their IRB of Record (all the time)?</th>
</tr>
</thead>
<tbody>
<tr>
<td>☐ Yes</td>
</tr>
<tr>
<td>☐ No</td>
</tr>
<tr>
<td>☐ It depends</td>
</tr>
</tbody>
</table>

Describe any board specialties of your IRB.

<table>
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<tr>
<th>_____________________________________________</th>
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<tr>
<th>Is your institution a covered entity?</th>
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</thead>
<tbody>
<tr>
<td>☐ Yes</td>
</tr>
<tr>
<td>☐ No</td>
</tr>
<tr>
<td>☐ Hybrid</td>
</tr>
</tbody>
</table>

Additional Comments

(Feel free to enter additional/clarifying info here)
Section 2: SITE-SPECIFIC LOCAL CONTEXT

This information may be helpful for a Reviewing IRB to understand when considering whether to serve as the Reviewing IRB for your organization.

Additional, study-specific information will be requested when a Reviewing IRB is reviewing a specific study for your site (e.g., COI, investigator training, study-specific consent requirements, state law or institutional requirements)

To what state laws is your institution subject

What circumstances affect age of consent in your state?
For example, in Pennsylvania a minor age 14 or above can consent to their own mental health treatment

Do you have any state or local laws or institutional policies that require RECORD KEEPING for longer than federal law requires under the Privacy Rule or Common Rule?

☐ Yes
☐ No

Please describe how long you are required to keep your records.

Do you require specific language in your consent form to describe what requires mandatory reporting to authorities?

☐ Yes, I will upload language in file
☐ Yes, I will insert language in text box
☐ No

Please upload the language required to be used around mandatory reporting to health authorities.

Please insert the language required to be used around mandatory reporting to health authorities.

Does your site require a site-specific logo appear on consent forms and/or recruitment documents?

☐ Yes
☐ No

Please upload the logo

Does your institution require approval of a waiver of authorization under HIPAA for review of medical records to identify eligible subjects?

☐ Yes
☐ No
If yes, the local context questions from section 2 are repeated for the component. Multiple component sites can be reported on in the site's Institutional Profile.
Local Considerations: Study-specific Human Research Protections Reviews (completed by the HRPP Liaison)
Study-Specific Local Context Worksheet

Please have this form completed by someone at your local institution who has knowledge of the protocol and any local regulatory, policy or community considerations that may impact the approvability of the research at your site.

Study Title

Local Site Name

STUDY TEAM INFORMATION

Local Site PI First Name

Local Site PI Last Name

Local Site PI Email

Please review the planned list of personnel who will be engaged in human subjects research.

Has all required training for the conduct of the research at your site been completed for each individual, including human subjects protections training, GCP training, and HIPAA training, as applicable?

☐ Yes
☐ No
☐ Our institution delegates this responsibility to the PI

NOTE: If all study personnel have not met the required training, please have them do so before submitting this application.

Additional Information regarding the verification that all training requirements are met:

((optional))

Are all involved individuals at the institution for this protocol credentialed and/or appropriately qualified and meet the institution's standards for eligibility to conduct research?

☐ Yes
☐ No

Please review the planned list of personnel who will be engaged in human subjects research and indicate whether COI applies:

☐ I have verified these personnel do not have any financial interests to disclose
☐ I have verified any relevant interests have been disclosed per my institutional policy and managed, as applicable
☐ Our institutional policy requires the researcher to disclose conflicts to another office/department and send the management plan to the Reviewing IRB
Where the conflict of interest management plan requires additional language in the consent form, please provide the language that is to be included in the consent form.

Where any financial conflicts of interest have been identified that require a management plan, please upload the relevant management plan.

Provide institutional point of contact for questions related to local management plan (this person should be someone in the office/entity who prepared the management plan).

(Provide name, email and phone number)

**STUDY-SPECIFIC LOCAL REQUIREMENTS**

Please review the protocol and template consent form and verify in the box below that there are sufficient resources available at your site to carry out the research as planned. If any changes are required to the study plan related to the resources available at your site, please outline the required changes below.

Please review the protocol and template consent form, as provided to you by your local study team, and identify areas where there are unique state, local or federal regulatory requirements that apply to this study and describe any steps that must be taken to adhere to these requirements.

For Example: legally authorized representatives, state laws regarding confidentiality of specific types of health information, emancipated minors.

Please Note: In the box below, please outline any specific changes needed to ensure adherence with the requirements you have identified that are applicable to the conduct of this study at your site. This may include changes to the consent form to include any language required based on state law requirements [e.g. reportability of test results for infectious diseases].

If your site has no unique regulatory requirements, enter "None" below.

Please review the protocol and template consent form, as provided to you by your local study team, and identify any institutional requirements (e.g., recruitment, data security, remuneration) that apply to this study and any steps that must be taken to adhere to these requirements.

Please Note: In the box below, please outline any specific changes needed to ensure adherence with the requirements you have identified that are applicable to the conduct of this study at your site. This may include changes to the consent form to include any language required based on local requirements [e.g. any specific local policy requirements related to consent for future use of biospecimens].

If your site has no specific changes, enter "None" below.
Please identify any ancillary reviews required at your site [e.g. radiation safety review, review for research with biospecimens, etc.] that will be required before this study may be initiated at your site.

If no ancillary reviews are required, please indicate "None" below.

<table>
<thead>
<tr>
<th>Based on the ancillary reviews listed above, are there any site-specific ancillary reviews that may impact the IRB review?</th>
</tr>
</thead>
<tbody>
<tr>
<td>○ Yes  ○ No</td>
</tr>
</tbody>
</table>

What is the current status of the review and approval by that ancillary committee?

Provide details here (i.e. outcome, anticipated date of review) or upload documentation below

Upload documentation here (i.e. outcome, anticipated date of review) or provide details in text box above

COMMUNITY CONSIDERATIONS

Please review the protocol and template consent and identify whether there are any special characteristics/concerns of your community of which the reviewing IRB should be aware for this specific study. Please also outline any steps that must be taken to address these concerns.

It is possible that the Reviewing IRB may have additional questions about your local community. Please include the best contact below for additional questions about local context information.

Local Contact First Name

Local Contact Last Name

Local Contact Email

Local Contact Phone
CONSENT FORMS: Please review the template consent form and, with consideration for any local site regulatory or institutional requirements, please ensure any local site required consent language is included before uploading the ICF(s) here.

<table>
<thead>
<tr>
<th>Question</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>Does your institution require approval of a waiver of authorization under HIPAA for review of medical records to identify eligible subjects?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Does your IRB require HIPAA authorization forms be a separate document from the Informed Consent Form?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NOTE: If you uploaded HIPAA authorization templates to your Institutional Profile (IP), they will be included in your local considerations packet for the Reviewing IRB. Check your IP to make sure the document(s) are up to date.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Does your site require a site-specific logo appear on consent forms and/or recruitment documents?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Note: the site specific logo uploaded to your Institutional Profile (IP) will be included in your local considerations packet for the Reviewing IRB. Check your IP to make sure the logo is up to date.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Does your institution require consent version date to be included on the consent document?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Does your institution require participant initials on each page?</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Please review the template consent form, and with consideration for any local regulatory or institutional requirements, provide site required subject injury language. This text should be study specific as it will be incorporated into the consent form by the CIRB.

Costs to participants language

This text will be incorporated into the consent form by the CIRB.

HIPAA Authorization Language

This text will be incorporated into the consent form by the CIRB.

Tissue Banking Language

This text will be incorporated into the consent form by the CIRB.
### Additional Local Considerations

If the Reviewing IRB sent additional study-specific questions about your local context/local considerations, please upload your answers to those questions here.

### Section IV - Signatures/Attestations

<table>
<thead>
<tr>
<th>Role</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>IRB Point of Contact First Name</td>
<td>________________________</td>
</tr>
<tr>
<td>IRB Point of Contact Last Name</td>
<td>________________________</td>
</tr>
<tr>
<td>IRB Point of Contact Title</td>
<td>________________________</td>
</tr>
<tr>
<td>IRB Point of Contact Phone</td>
<td>________________________</td>
</tr>
<tr>
<td>IRB Point of Contact Email</td>
<td>________________________</td>
</tr>
</tbody>
</table>

By signing below, I attest that the information fulfills the relying institution's responsibilities, as outlined in the reliance agreement, for the provision of local context information and to the accuracy and completeness of the information provided herein.

__________________________________________

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**Confidential**
The PI Survey regarding Conduct of the Study at their Site
Local Context PI Survey

Study Title

Participating Site Name:

STUDY TEAM INFORMATION

Site Investigator’s Name

Site Investigator’s Email Address

Site Investigator’s Phone

(Consent Form contact information)

Lead Study Contact Name

Lead Study Contact Email

Lead Study Contact Phone

Please review the planned list of personnel who will be engaged in human subjects research.

Has all required training for the conduct of the research at your site been completed for each individual, including human subjects protections training, GCP training, and HIPAA training, as applicable?

☐ Yes
☐ No
☐ Our institution delegates this responsibility to the HRPP

NOTE: If all study personnel have not met the required training, please have them do so before submitting this application.

Additional information regarding the verification that all training requirements are met.

((optional))

Please review the planned list of personnel who will be engaged in human subjects research and indicate whether COI applies:

☐ I have verified there are no financial interests to disclose
☐ I have verified any relevant interests have been disclosed per my institutional policy and managed, as applicable
Where the conflict of interest management plan requires additional language in the consent form, please provide the language that is to be included in the consent form.

Where any financial conflicts of interest have been identified that require a management plan, please upload the relevant management plan.

Provide institutional point of contact for questions related to local management plan (this person should be someone in the office/entity who prepared the management plan).

CONSENT DOCUMENT

Person to Contact

This text will be incorporated into the consent form by the CIRB.

Is there any additional language about how the study will be conducted at your local site that needs to be disclosed on the consent document?

☐ Yes
☐ No

Please provide language to include under the following subheading of the consent document - Other information about how the study will be done at your local site:

This text should be study specific as it will be incorporated into the consent form by the CIRB.

SITE SPECIFIC PLANS FOR RECRUITMENT, CONSENTING, AND DATA AND SAFETY MONITORING

RECRUITMENT PLAN: Are there any differences to the initial contact and/or recruitment plan at your site from that described in the protocol or associated documents based on local requirements or state law?

☐ Yes
☐ No

How does the recruitment plan differ?

Please describe the specific steps to be used to identify and/or contact prospective participants at your site. Also, if applicable, describe how you have access to lists of potential participants.
CONSENTING PLAN: Does the consent plan for your site differ in any way from that outlined in the protocol?

- Yes
- No

How does the consent plan differ?

Please describe the specific steps for obtaining informed consent and the procedures that will be utilized to protect the privacy of individuals.

Does the person obtaining consent have an existing relationship with the participant(s)?

- Yes
- No

Please indicate/describe the relationship(s) and how you will protect against undue influence or coercion.

DATA AND SAFETY MONITORING PLAN:

Does the data and safety monitoring plan for your site differ in any way from that outlined in the protocol?

- Yes
- No

How does the data and safety monitoring plan differ?

Describe plans for monitoring the progress of trials and safety of participants at your site.

Are there any other different requirements for how the protocol will be implemented and/or conducted at your site based on local requirements or state laws?

- Yes
- No

Please describe any other different requirements for how the protocol will be implemented at your site.
SIGNATURES/ATTESTATIONS

By signing below, I attest to the accuracy and completeness of the information provided herein.

________________________________________