

Communication Plan

COMMUNICATION RESPONSIBILITY	RESPONSIBLE PARTY	NOTES
<p>COI: Providing applicable conflict of interest management plans for relying site study teams to the Reviewing IRB</p>	<p><input type="checkbox"/> Reviewing IRB <input type="checkbox"/> Lead Study Team <input type="checkbox"/> Relying Site Study Team(s) <input type="checkbox"/> Relying Site(s) POC(s) <input type="checkbox"/> Other, specify:</p>	
<p>STUDY TEAM TRAINING & QUALIFICATIONS: Providing confirmation to the Reviewing IRB that relying site study teams have completed relevant training and are qualified to conduct the proposed research</p>	<p><input type="checkbox"/> Reviewing IRB <input type="checkbox"/> Lead Study Team <input type="checkbox"/> Relying Site Study Team(s) <input type="checkbox"/> Relying Site(s) POC(s) <input type="checkbox"/> Other, specify:</p>	
<p>LOCAL CONTEXT INFORMATION: Providing local context information to the Reviewing IRB regarding state laws and institutional requirements that pertain to the review of the ceded study</p>	<p><input type="checkbox"/> Reviewing IRB <input type="checkbox"/> Lead Study Team <input type="checkbox"/> Relying Site Study Team(s) <input type="checkbox"/> Relying Site(s) POC(s) <input type="checkbox"/> Other, specify:</p>	
<p>IRB APPLICATION – STUDYWIDE: Preparing and submitting the study-wide application for initial IRB review and study-wide amendments to the Reviewing IRB</p>	<p><input type="checkbox"/> Reviewing IRB <input type="checkbox"/> Lead Study Team <input type="checkbox"/> Relying Site Study Team(s) <input type="checkbox"/> Relying Site(s) POC(s) <input type="checkbox"/> Other, specify:</p>	

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<p>IRB APPLICATION – SITE-SPECIFIC: Preparing and submitting the site-specific applications and site-specific amendments to the Reviewing IRB that address site variations in study conduct, informed consent language, HIPAA Privacy Rule requirements, subject identification and recruitment processes (including recruitment materials), and any other applicable components of the research</p>	<ul style="list-style-type: none"> <input type="checkbox"/> Reviewing IRB <input type="checkbox"/> Lead Study Team <input type="checkbox"/> Relying Site Study Team(s) <input type="checkbox"/> Relying Site(s) POC(s) <input type="checkbox"/> Other, specify: 	
<p>IRB DETERMINATIONS: Providing documentation of IRB determinations to relying site study teams</p>	<ul style="list-style-type: none"> <input type="checkbox"/> Reviewing IRB <input type="checkbox"/> Lead Study Team <input type="checkbox"/> Relying Site Study Team(s) <input type="checkbox"/> Relying Site(s) POC(s) <input type="checkbox"/> Other, specify: 	
<p>IRB-APPROVED DOCUMENTS: Providing copies of IRB-approved materials to the lead study team</p>	<ul style="list-style-type: none"> <input type="checkbox"/> Reviewing IRB <input type="checkbox"/> Lead Study Team <input type="checkbox"/> Relying Site Study Team(s) <input type="checkbox"/> Relying Site(s) POC(s) <input type="checkbox"/> Other, specify: 	
<p>IRB-APPROVED DOCUMENTS – RELYING SITES: Providing copies of the most current versions of IRB-approved materials to relying site study teams in a timely manner</p>	<ul style="list-style-type: none"> <input type="checkbox"/> Reviewing IRB <input type="checkbox"/> Lead Study Team <input type="checkbox"/> Relying Site Study Team(s) <input type="checkbox"/> Relying Site(s) POC(s) <input type="checkbox"/> Other, specify: 	

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<p>CONSENT FORM TEMPLATE: Providing the consent form template to relying site study teams</p>	<ul style="list-style-type: none"> <input type="checkbox"/> Reviewing IRB <input type="checkbox"/> Lead Study Team <input type="checkbox"/> Relying Site Study Team(s) <input type="checkbox"/> Relying Site(s) POC(s) <input type="checkbox"/> Other, specify: 	
<p>CONSENT FORM LANGUAGE: Incorporating site-specific language into consent form(s) and providing these consent form(s) to the Reviewing IRB</p>	<ul style="list-style-type: none"> <input type="checkbox"/> Reviewing IRB <input type="checkbox"/> Lead Study Team <input type="checkbox"/> Relying Site Study Team(s) <input type="checkbox"/> Relying Site(s) POC(s) <input type="checkbox"/> Other, specify: 	
<p>REVIEWING IRB POLICIES: Providing relevant Reviewing IRB policies to the lead study team</p>	<ul style="list-style-type: none"> <input type="checkbox"/> Reviewing IRB <input type="checkbox"/> Lead Study Team <input type="checkbox"/> Relying Site Study Team(s) <input type="checkbox"/> Relying Site(s) POC(s) <input type="checkbox"/> Other, specify: 	
<p>CONTINUING REVIEW INFORMATION: Obtaining and collating study-wide information for continuing review to the Reviewing IRB</p>	<ul style="list-style-type: none"> <input type="checkbox"/> Reviewing IRB <input type="checkbox"/> Lead Study Team <input type="checkbox"/> Relying Site Study Team(s) <input type="checkbox"/> Relying Site(s) POC(s) <input type="checkbox"/> Other, specify: 	

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<p>CONTINUING REVIEW SUBMISSION: Submitting continuing review progress report to the Reviewing IRB</p>	<p>Reviewing IRB Lead Study Team Relying Site Study Team(s) Relying Site(s) POC(s) Other, specify:</p>	
<p>REPORTABLE EVENTS: Reporting reportable events to the Reviewing IRB (e.g., unanticipated problems, noncompliance, subject complaints)</p>	<p>Reviewing IRB Lead Study Team Relying Site Study Team(s) Relying Site(s) POC(s) Other, specify:</p>	
<p>CLOSURE REPORTS: Providing the Reviewing IRB with required information when a study is closed.</p>	<p>Reviewing IRB Lead Study Team Relying Site Study Team(s) Relying Site(s) POC(s) Other, specify:</p>	
<p>OTHERS – CONSIDER IF APPLICABLE</p>		
<p>RADIATION SAFETY</p>		
<p>INSTITUTIONAL BIOSAFETY COMMITTEE</p>		
<p>LABORATORY SAFETY</p>		