

# File Checklist for Human Subject/Clinical Trial Form



Needed for each study related to the project. This information is now located on new Human Subject & Clinical Trials Information form. *This form is a mixture of uploads and entered data, (sections 1.1, 2.1& 2.2, 4.1, 4.2a&c, 4.3 among others).* **Questions 1.4 A-D will determine if your project is considered a Clinical Trial, if you answer no to just one of these four questions your project is not considered a Clinical Trial.** If you considered a use of Human Subject project and not deemed a clinical trial, you only have to complete sections 1-3, Clinical Trial proposals also have to complete sections 4 & 5.

File Type	Comments	Page Limits
Item 2.4: Inclusion of Women, Minorities and Children	These are now all uploaded together. <b>As of 1/25/2019 you need to include Inclusion Across Lifespan</b> ( <a href="https://grants.nih.gov/grants/funding/lifespan/lifespan.htm">https://grants.nih.gov/grants/funding/lifespan/lifespan.htm</a> )	No limit
Item 2.5: Recruitment and Retention Plan	This is new with the E Forms	No limit
Item 2.7: Study Timeline	This is new with the E Forms	No limit
Item 3.1: Protection of Human Subjects*		No limit
Item 3.2: Single IRB Plan	New Mandatory if this is a domestic multi-site study	No limit
Item 3.3: Data & Safety Monitoring Plan	Mandatory for CT projects, optional for HS only submissions	No limit
Item 3.5: Overall structure of the study team	Optional	No limit
Item 4.4: Statistical Design and Power	Mandatory for CT projects unless otherwise noted in the FA	No limit
Item 4.5: Subject Participation Duration	Mandatory for CT projects unless otherwise noted in the FA	No limit
Item 4.7: Dissemination Plan	Mandatory for CT projects unless otherwise noted in the FA	No limit

	Data Entry Fields	Comments	Character Limit/Notes
	Item 2.1: Conditions or Focus of Study	A list of keywords (search criteria) that describe the study	None
	Item 2.2: Eligibility Criteria	Inclusion and exclusion conditions	15,000 Characters
	Item 2.3: Age Limits (Minimum and Maximum)	Need to include age as one of the options from a drop down that include: years, months, weeks, days, hours, minutes	
	Item 2.6: Recruitment Status	Enter or select from a drop down	Choices include but are not limited to: not yet recruiting, recruiting, completed, enrolling by invitation and active
	Item 2.8: Enrollment of first Subject	You need to enter both a date and whether that is an actual or anticipated target	
	Inclusion Enrollment Report	This needs to be activated and completed	If doing a secondary data analysis figure need to be entered in the actual and not planned section
	Item 4.1: Brief Summary	Description and objectives of the protocol including the primary and secondary endpoints.	Limited to 5,000 Characters
	Item 4.2a: Narrative of Study Description	Describe your plans for assignment of participants and delivery of interventions. You will also need to show that your methods for sample size and data analysis are appropriate given those plans. For trials that randomize groups or deliver interventions to groups, special methods are required.	Limited to 32,000 Characters
	Item 4.2b: Primary Purpose	Enter or select from the dropdown menu a single "Primary Purpose" that best describes the clinical trial	Choices include: Treatment, Prevention, Diagnostics, Screening, Basic Science, and Device Feasibility
	Item 4.2c: Interventions	Complete fields for each intervention to be used in your proposed protocol. If an arm of the study to which subjects will be assigned (as discussed in 4.2.a. Narrative Study Description) includes more than one intervention (e.g., drug plus educational intervention), complete this section for each intervention.	You can add up to 20 interventions. You must choose an intervention, and a unique name, limited to 200 characters, and enter a description of up to 1,000 characters
	Item 4.2d: Study Phase	Enter or choose from a drop down menu	Choices range from Early Phase 1 (or Phase 0) to Phase 4
	Item 4.2e: Intervention Model	Enter or choose from a drop down menu	Choices include: Single Group, Parallel, Factorial, and Sequential
	Item 4.3 Outcome Measures - Primary, Secondary and other important measures	Needs to be completed for each outcome: <ul style="list-style-type: none"> <li>primary (the outcome measures specified in your protocol that are of greatest importance to your study),</li> <li>secondary (measures specified in your protocol that are of lesser</li> </ul>	Note: You may have more than one primary outcome measure, and you can add up to 50 unique outcome measures

		<p>importance to your study than your primary outcomes), and other important measures (additional key outcome measures used to evaluate the intervention to be collected during your proposed clinical trial.).</p> <ul style="list-style-type: none"> <li>• other important measures (additional key outcome measures used to evaluate the intervention to be collected during your proposed clinical trial.).</li> </ul> <ol style="list-style-type: none"> <li>1. Time Frame: Indicate when a measure will be collected for analysis (e.g., baseline, post-treatment).</li> <li>2. Brief Description: Describe the metric used to characterize the outcome measure if the metric is not already included in the outcome measure name.</li> <li>3. Your description is additional key outcome measures used to evaluate the intervention.</li> </ol>	<p>The Description is limited to limited to 999 characters.</p> <p>The Time Frame and Outcome Measure is limited to 255 Characters</p>
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**Note:** You should always read the specific FA to see if it deviates from the standard requirements. You should also see if any other files or appendix material are allow or required. If it is not specified these items should not be included. Contact The Office of Clinical Award Administration 8-4490/[SOMCAA@stonybrookmedicine.edu](mailto:SOMCAA@stonybrookmedicine.edu), if you have any questions on what is required for a complete submission.