

## **ClinicalTrials.gov Registration Designation Letter**

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(the "Trial") has been identified as a study for which registration is required on					
ClinicalTrials.gov. This document serves as notice that	has				
designated the Principal Investigator,	, of the Trial as the				
"Responsible Party" for the purposes of registering the Trial with	the ClinicalTrials.gov registry.				
As the Responsible Party, you are responsible for accurately registering this clinical trial on					
ClinicalTrials.gov, as required by the Food and Drug Administration Amendments Act					
(FDAAA).					

To assist in fulfilling responsibilities to register on ClinicalTrials.gov, Partners Human Research Affairs has made the following resources available:

Partners Clinical Trials Registration Information <a href="http://healthcare.partners.org/phsirb/investigatorctregistration.htm">http://healthcare.partners.org/phsirb/investigatorctregistration.htm</a>

Generally speaking, requirements include:

- □ Registration of the study on ClinicalTrials.gov.
- □ Updates every 6 months as required by ClinicalTrials.gov.
- □ Notification to ClinicalTrials.gov within 30 days of study completion.

My signature below acknowledges that I, as the "Responsible Party" by following statutory requirements:

have been designated and assures that I meet the

- 1) I am responsible for conducting the Trial.
- 2) I have access to and control over the data from the Trial.
- 3) I have the right to publish the results of the Trial.
- 4) I have the ability to meet all of the requirements under FDAAA for the submission of clinical trial information.

PI Signature:	 Date: