

Office of Sponsor and Regulatory Oversight

Protocol Signature Page

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Instructions

This form should be completed and signed by the local Principal Investigator who is responsible for the study implementation at his/her specific site (i.e., if the study is under an IND this page is signed by the individual who signs the Form FDA 1572; or, if the study is under an IDE this page is signed by the individual who signs the Investigator of Record form).

Provide the following info	ation and a digital or wet signature. A reproduced signature will not be accepte	d.
Protocol Number:		
Protocol Version:		
Protocol Date:		
Site Name:		
of the protocol, including	the necessary assurance that this trial will be conducted according to all stipul statements regarding confidentiality, and according to local legal and regulator US federal regulations and ICH E6 Good Clinical Practice (GCP) guidelines.	
I agree to conduct the stu	n compliance with GCP and applicable regulatory requirements.	
-	in accordance with the current protocol and will not make changes to the prond IRB approval, except when necessary to protect the safety, rights, or well	
Principal Investigator (pri	ype):	
Signed:	Date:	

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