# How to Create a Study CTMS version 2.1 APPLICABLE TO: Company Admin Study Manager CRA Note: In order to successfully create a study, the sponsor and product must be created first. I. Log into the CTMS and click on Studies on the left.

2 To add a new study, click Add.



Enter the study details, then click
 Create to save.

Create Study		□ <b>x</b>
Sponsor*		
		~
Protocol Number*	Study Name*	
		0
Protocol Title*		
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Types o Types		
Select		
Select		
Select	Status Date*	





December 2024 Page 1 of 2

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**Note**: At this point, the study itself has been created. The following are recommendations for additional steps in order to complete the setup of the study in the CTMS.



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# Navigate to the Study Details area to manage Study Settings.

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		4 © Settings

Navigate to the **Access Permissions** area and indicate the members of your study team and their study-level access level.

- All non-CSM users who will play a part in the study should be added with a viewonly level of access.
- If the study is blinded, you will need to reach out to the Trial Interactive Service Desk at <u>help@trialinteractive.com</u> to have an Unblinded CSM created so that that user can manage unblinded aspects of the study including designating unblinded CRAs.
- For additional information, please see the related job aid and video regarding how to manage study access permissions.

Access Permissions						
0 4	dd 🖪 Edit Permissions ඟ A		S Remove Set as Owner			
Теа	m Members Only All with Access	Sta	tus All ~ Access Permission All ~			
2 Use	ers 1 Selected					
	Name	1	Status	Access Permission		
~	🐣 CRA 003		ACTIVE	Study Viewer		
	CSM 003 Owner		ACTIVE	Unblinded - CSM		



## In the **Settings** Area:

- If applicable to your study, indicate which users are to be tasked with reviewing site visit reports. This list comes from those with access to the study. Please note that Blinded and Unblinded Reviewers need to be indicated separately, even if the same people will be performing both functions.
  - If this is not done, the Visit Report workflow will not function as expected, leading to delays and potential document rejections.

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Sponsor Diagnova Therapeutics Product Diagnovate	Reviewers				
🔅 General		Q. Search		Q Search	
Regions	1 Blinded Reviewer		1 Unblinded Reviewers		
Milestone Templates	Name	Email	Name	Email	
	CSM 003	CSM003@ti.com	CSM 003	CSM003@ti.com	



December 2024 Page 2 of 2