The YODA Project Research Proposal Due Diligence Assessment

Part 1: General Information			
YODA Project (Protocol) ID:		2020-4215	
Date:		18 March 2020	
Product Name:		Bosentan/Macitentan	
Therapeutic Area:		Pulmonary Hypertension	
Product Class:		Endothelin receptor antagonist	
Condition(s) Studied:		Idiopathic Pulmonary Fibrosis	
Protocol Number(s) and Title(s):		 NCT00391443 AC-052-321 Effects of Bosentan on Morbidity and Mortality in Patients With Idiopathic Pulmonary Fibrosis - a Multicenter, Double-blind, Randomized, Placebo-controlled, Parallel Group, Eventdriven, Group Sequential, Phase III Study NCT00071461 AC-052-320 (BUILD 1) - A Double-blind, Randomized, Placebo- controlled, Multicenter Study to Assess the Efficacy, Safety, and Tolerability of Bosentan in Patients With Idiopathic Pulmonary Fibrosis, Open Label Extension NCT0090331 AC-055B201 A Double-blind, Randomized, Placebo-controlled, Multicenter, Parallel Group Study to Evaluate the Efficacy, Safety, and Tolerability of Macitentan in Patients With Idiopathic Pulmonary Fibrosis 	
		Part 2: Data Availability	,
Question:			Response:
Data Holder has authority to provide clinical trial data or development partner has agreed to s clinical trial data.			Yes
	/A o olootropio oli	nical trial data or data can be converted to electronic format.	Vac
	A electronic cli	nical that data of data call be converted to electronic format.	Yes
,		cal trial data in accordance with current HIPAA and EU criteria	Yes
allows protection of participant privacy and confidentiality.			
Comments: N/A			
The product and relevan	t indication st	udied has either been approved by regulators in the US and EU,	Yes
or terminated from development.			
Comments: N/A			
•		trial and trial has been completed for a period of at least 18	Yes
months (or results published in peer-reviewed biomedical literature). Comments: N/A			
Comments: N,			
	F	Part 3: Data Availability Summary	
Based on the responses to the above Data Availability questions, the requested clinical trial data can			Yes
be made available for data sharing.			
		Part 4: Proposal Review	
Question:			Response:
Summary-level CSR data is appropriate for the proposed analysis.			No
Participant-level data is appropriate for the proposed analysis.			Yes
	erway or comp	leted/pending disclosure by Janssen.	No
Comments:			