The YODA Project Research Proposal Due Diligence Assessment

Part 1: General Information		
YODA Project (Protocol) ID:	2016-0774	
Date:	21 March 2016	
Product Name: PROCRIT		
Therapeutic Area:	ONC - EP	
Product Class:	Hematologic Agents	
Condition(s) Studied: Anemia		
Protocol Number(s) and Title(s):	NCT00091910 - A Randomized, Double-Blind, Placebo-Controlled Study to Determine the Efficacy and Safety of Epoetin Alfa in Critically III Subjects EPO-2 /// PR98-15-014-No NCT number- Efficacy in the rHuEPO (Epoetin Alfa) in the Critically III Patient: A Randomized, Double Blind, Placebo-Controlled trial	
Part 2: Data Availability		
	Question:	Response:
Data Holder has authority to p partner has agreed to share cl Comments: N/A	rovide clinical trial data or development	Yes
Data Holder has sharable electronic clinical trial data or data can be converted to electronic format.		Yes
Comments: N/A De-identification and redaction of clinical trial data in accordance with current HIPAA and EU criteria allows protection of participant privacy and confidentiality.		Yes
Comments: N/A		
The product and relevant indication studied has either been approved by regulators in the US and EU, or terminated from development. Comments: N/A		Yes
Comments: N/A Data Holder has completed the clinical trial and trial has been completed for a period of at least 18 months (or results published in peer-reviewed biomedical literature). Comments: N/A		
<u> </u>	Part 2: Data Availability Summary	
Part 3: Data Availability Summary Based on the responses to the above Data Availability questions, the Yes		
requested clinical trial data can be made available for data sharing.		Yes
Part 4: Proposal Review		
Question:		Response:
Summary-level CSR data is appropriate for the proposed analysis.		Yes
Participant-level data is appropriate for the proposed analysis.		Yes
A similar analysis is underway or completed/pending disclosure by Janssen.		No
Comments:		