The YODA Project Research Proposal Due Diligence Assessment

Part 1: General Information			
YODA Project (Protocol) ID:	2022-5037		
Date:	22 September 2022-Updated 4Oct22		
Product Name:	Infliximab/Golimumab/Ustekinumab		
Therapeutic Area:	Immunology		
Product Class:	Antirheumatic Agents - Biologic Response Modifiers		
Condition(s) Studied:	Ulcerative Colitis		
Protocol Number(s) and Title(s):	NCT00036439 - C0168T37 - A Randomized, Placebo-controlled, Double-blind Trial to Evaluate the Safety and Efficacy of Infliximab in Patients With Active Ulcerative Colitis NCT02407236 - CNT01275UC03001 - A Phase 3, Randomized, Double-blind, Placebo-controlled, Parallel group, Multicenter Protocol to Evaluate the Safety and Efficacy of Ustekinumab NCT00488631 - C0524T18 - A Phase 3 Multicenter, Randomized, Placebo-controlled, Double-blind Study to Evaluate the Safety and Efficacy of Golimumab Maintenance Therapy, Administered Subcutaneously, in Subjects With Moderately to Severely Active Ulcerative Colitis NCT00096655 - C0168T46 - A Randomized, Placebo-controlled, Double-blind Trial to Evaluate the Safety and Efficacy of Infliximab in Patients With Active Ulcerative Colitis		
Part 2: Data Availability			
Data Holder has authority to provide clinical trial data or development partner has agreed to share clinical trial data. Comments: Data Holder has sharable electronic clinical trial data or data can be converted Yes		Yes	
to electronic format.			
Comments: De-identification and redaction of clinical trial data in accordance with current HIPAA and EU criteria allows protection of participant privacy and confidentiality.			
Comments: The product and relevant indication studied has either been approved by regulators in the US and EU, or terminated from development. Yes			
Comments: 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:			
Part 3: Data Availability Summary			
Based on the responses to the above Data Availability questions, the requested clinical trial data are available for a data sharing request.			

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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		
A similar analysis is underway or completed/pending disclosure by Janssen.	No		
Comments:			