The YODA Project Research Proposal Due Diligence Assessment

Part 1: General Information			
YODA Project (Protocol) ID:	2017-1676		
Date:	16 May 2017		
Product Name:	Paliperidone palmitate		
Therapeutic Area:	Neuroscience		
Product Class:	atypical antipsychotics		
Condition(s) Studied:	Schizophrenia		
Protocol Number(s) and Title(s):	NCT00518323- A Randomized, Multicenter, Double-Blind, Weight-Based, Fixed-Dose, Parallel-Group, Placebo-Controlled Study of the Efficacy and Safety of Extended Release Paliperidone for the Treatment of Schizophrenia in Adolescent Subjects, 12 to 17 Years of Age NCT00334126- A Randomized, Double-blind, Placebo-controlled, Parallel Group Study to Evaluate the Efficacy and Safety of Paliperidone ER Compared to Quetiapine in Subjects With an Acute Exacerbation of Schizophrenia NCT00085748- A Randomized, 6-Week Double-Blind, Placebo-Controlled Study With an Optional 24-Week Open-Label Extension to Evaluate the Safety and Tolerability of Flexible Doses of Paliperidone Extended Release in the Treatment of Geriatric Patients With Schizophrenia NCT00078039- Trial Evaluating Three Fixed Dosages of Paliperidone Extended-Release (ER) Tablets and Olanzapine in the Treatment of Patients With Schizophrenia NCT00077714- A Randomized, Double-blind, Placebo- and Active-controlled, Parallel-group, Dose-response Study to Evaluate the Efficacy and Safety of 2 Fixed Dosages of Paliperidone Extended Release Tablets and Olanzapine, With Open-label Extension, in the Treatment of Patients With Schizophrenia NCT00083668- A Randomized, Double-blind, Placebo- and Active-controlled, Parallel-group, Dose-response Study to Evaluate the Efficacy and Safety of 3 Fixed Dosages of Paliperidone Extended Release (ER) Tablets and Olanzapine, With Open-label Extension, in the Treatment of Patients With Schizophrenia NCT00524043- A Randomized, Double-Blind, Placebo- and Active-Controlled, Parallel-Group Study to Evaluate the Efficacy and Safety of a Fixed Dosage of 1.5 mg/Day of Paliperidone Extended Release (ER) in the Treatment of Subjects With Schizophrenia		

The YODA Project Research Proposal Due Diligence Assessment

Question: Data Holder has authority to provide clinical trial data or development partner has agreed to share clinical trial data. Comments: N/A Data Holder has sharable electronic clinical trial data or data can be converted to electronic format. 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. 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 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 Part 3: Data Availability Summary Based on the responses to the above Data Availability questions, the requested clinical trial data can be made available for data sharing.	Response: Yes
Part 3: Data Availability Summary Data Holder has sharable electronic clinical trial data or data can be converted to electronic format. 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. 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 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 Part 3: Data Availability Summary Based on the responses to the above Data Availability questions, the	
Data Holder has sharable electronic clinical trial data or data can be converted to electronic format. 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. 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 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 Part 3: Data Availability Summary Based on the responses to the above Data Availability questions, the	Yes
to electronic format. 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. 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 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 Part 3: Data Availability Summary Based on the responses to the above Data Availability questions, the	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. 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 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 Part 3: Data Availability Summary Based on the responses to the above Data Availability questions, the	
De-identification and redaction of clinical trial data in accordance with current HIPAA and EU criteria allows protection of participant privacy and confidentiality. 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 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 Part 3: Data Availability Summary Based on the responses to the above Data Availability questions, the	
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 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 Part 3: Data Availability Summary Based on the responses to the above Data Availability questions, the	Yes
regulators in the US and EU, or terminated from development. 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 Part 3: Data Availability Summary Based on the responses to the above Data Availability questions, the	
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 Part 3: Data Availability Summary Based on the responses to the above Data Availability questions, the	Yes
period of at least 18 months (or results published in peer-reviewed biomedical literature). Comments: N/A Part 3: Data Availability Summary Based on the responses to the above Data Availability questions, the	
Part 3: Data Availability Summary Based on the responses to the above Data Availability questions, the	Yes
Based on the responses to the above Data Availability questions, the	
	Yes
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. Comments:	No

Part 1: General Information			
YODA Project (Protocol) ID:	2017-1676		
Date:	16 May 2017		
Product Name:	Risperidone		
Therapeutic Area:	Neuroscience		
Product Class:	atypical antipsychotics		
Condition(s) Studied:	Schizophrenia		

The YODA Project Research Proposal Due Diligence Assessment

Protocol Number(s) and Title(s):	· · · · · · · · · · · · · · · · · · ·			
	NCT00088075- A Randomized, Double-Blind, Pla Clinical Study of the Efficacy and Safety of Rispe Treatment of Schizophrenia in Adolescents			
	NCT# N/A- RIS-USA-1- Risperidone versus haloge in the treatment of schizophrenia	peridol versus placebo		
	NCT# N/A- RIS-USA-72- The safety and efficacy qd and 4 mg qd compared to placebo in the tre schizophrenia			
	Part 2: Data Availability			
	Question:	Response:		
Data Holder has authority to provide clinical trial data or development partner has agreed to share clinical trial data.		Yes		
Comments: N/A				
Data Holder has sharable electronic clinical trial data or data can be converted		Yes		
to electronic format.				
Comments: N/A		Г		
De-identification and redaction HIPAA and EU criteria allows pure confidentiality.	Yes			
Comments: N/A				
The product and relevant indic	Yes			
regulators in the US and EU, or terminated from development.				
Comments: N/A				
Data Holder has completed the period of at least 18 months (o biomedical literature).	Yes			
Comments: N/A				
P	art 3: Data Availability Summary			
Based on the responses to the	Yes			
requested clinical trial data car				
	D. 14 D 12			
Part 4: Proposal Review				
Question:		Response:		
Summary-level CSR data is appropriate for the proposed analysis.		Yes		
Participant-level data is approp	Yes			
A similar analysis is underway of Comments:	or completed/pending disclosure by Janssen.	No		