

## Non Serious Adverse Drug Reactions Report

Start Date:2016-06-17 End Date:2016-06-20

Study :EMR700623-541		Investigator :Ying Zhong		Country of Investigator :China		SiteNo:C05	
Subject No :C05-0021		Subject Initials :FYZ		DOB :08/28/1990		Sex:Female	
				Race:Asian		Height:150cm	
						Weight:45kg	
First administration date of batch :				Batch number :			
Study Drug		Start Date		Dose		Change in Dose	
Gonal-f New Pen Stimulation Treatment		08/04/2015		225			
Adverse Event		Start Date		End Date		Time related to study treatment	
OHSS		08/18/2015		08/21/2015		Causality to study drug	
						Severity	
						Related	
						Mild	
Causality Factors		Action Taken with Study Treatment		Other action taken		Outcome	
Protocol procedure**, Concomitant medication**(Other value:)		Not applicable		None		Resolved	
Event Description:abdominal distention;Drug : Hydroxyethyl Starch 130/0.4 and Sodium Chloride Injection 500ml , ivgtt bid calcium gluconate injection 10ml+dextrose injection 500ml ivgtt qd 19-Aug-2015 / 21-Aug-2015							
Subject received concomitant medications:							
Does the subject have any relevant past or present medical conditions:Yes							
Condition				Start Date		Related to study condition	
salpingocatheterism				UK-Unk-2012		Ongoing	
						Not on treatment/medication	

18-JUN-16

Ying  
Zhong/EMR700623-541/C05-0021

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Start Date:2016-06-17 End Date:2016-06-20

Study :EMR700623-541	Investigator :Ying Zhong		Country of Investigator :China	SiteNo:C05		
Subject No :C05-0041	Subject Initials :W-Y	DOB :07/03/1981	Sex:Female	Race:Asian	Height:153cm	Weight:55kg
First administration date of batch :			Batch number :			
Study Drug	Start Date	Dose	Change in Dose			
Gonal-f New Pen Stimulation Treatment	08/08/2015	150				
Adverse Event	Start Date	End Date	Time related to study treatment	Causality to study drug	Severity	
OHSS	08/23/2015	08/25/2015		Related	Mild	
Causality Factors		Action Taken with Study Treatment	Other action taken	Outcome	AE Special Interest	AE dose limiting toxicity
Protocol procedure**,Concomitant medication**(Othervalue:)		Dose not changed	Led to study termination	Resolved		
Event Description:abdominal distention;Drug : Hydroxyethyl Starch 130/0.4 and Sodium Chloride Injection 500ml , ivgtt bid calcium gluconate injection 10ml+dextrose injection 500ml ivgtt qd 21-Aug-2015 / 25-Aug-2015						
Subject received concomitant medications:						
Does the subject have any relevant past or present medical conditions:No						
Condition				Start Date	Related to study condition	Ongoing

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Study :EMR700623-541	Investigator :Ying Zhong		Country of Investigator :China	SiteNo:C05		
Subject No :C05-0132	Subject Initials :HQL	DOB :07/16/1980	Sex:Female	Race:Asian	Height:166cm	Weight:57kg
First administration date of batch :			Batch number :			
Study Drug	Start Date	Dose	Change in Dose			
Gonal-f New Pen Stimulation Treatment	08/27/2015	225				
Adverse Event	Start Date	End Date	Time related to study treatment	Causality to study drug	Severity	
OHSS	09/09/2015	09/16/2015		Related	Mild	
Causality Factors		Action Taken with Study Treatment	Other action taken	Outcome	AE Special Interest	AE dose limiting toxicity
Disease under study**(Othervalue:)		Dose not changed	None	Resolved		
Event Description:						
Subject received concomitant medications:						
Does the subject have any relevant past or present medical conditions:Yes						
Condition				Start Date	Related to study condition	Ongoing
Fallopian tube repair anaplasty				UK-Unk-2011	Not on treatment/medicatio n	

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