Non Serious Adverse Drug Reactions Report  Start Date: 2015-10-21 End Date: 2015-10-22								
Subject No :C02-0111	Subject Initials :CXW	<b>DOB</b> :10/10/1979	Sex:Female	Race: Asian	Height:162cm	Weight:68.0kg		
First administration date of batch :			Batch number :	number:				
Study Drug	Start Date		Dose	Change in Dose	•			
Gonal-f New Pen Stimulation Treatment	05/20/2015		200					
Adverse Event	Start Date	End Date	Time related to study treatment	Causality to study drug	Severity			
OHSS				Related	Moderate			
Causality Factors		Action Taken with Study Treatment	Other action taken	Outcome	AE Special Interest	AE dose limiting toxicity		
None(Othervalue:)		Not applicable	Led to study termination	Resolved				
Event Description:Moderate OHSS patients, improved canceled after embryo transfer.								
Subject received concor	mitant medications:							
Does the subject have any relevant past or present medical conditions: Yes								
Condition				Start Date	Related to study condition	Ongoing		
Ectopic pregnancy laparoscopic surgery, the left fallopian tube embryo window.				Uk-Unk-2012	Not on treatment/medication			
Tubal lipiodol angiography				Uk-Unk-2013	Not on treatment/medication			

21-OCT-15 NA/EMR700623-541/C02-0111