Individual Case Safety Report Form

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General Information								
EudraVigilance Local Report Number EU			EU-EC-10011778850					
Sender Type			Regulatory authority					
Sender's Organisation			EEA Regulator					
Type of Report			Spontaneous					
Primary source country			European Economic Area					
Reporter's qualification			Non-Healthcare Professional					
Case serious?			Yes					
Patient								
Age Group			Age Group (as per reporter)			Sex		
18-64 Years						Female		
Reaction	on / Event							
MedDRA LLT			Duration	Outcome		•	Seriousness ¹	
Foetal malpresentation			78.0 Days	Unknown			other	
Drug In	formation							
Role ²	Drug		Duration	Dose	U	Inits in Interval	Action taken	
S	COMIRNATY - TOZINAMERAN			1.0 (DF)				
Drug Information (cont.)								
Info ³	Drug	Indication		Pha		arm. Form	Route of Admin.	
	COMIRNATY - TOZINAMERAN		COVID-19 immunisation					

Seriousness: death=results in death; life threat.=life threatening; hospital.=requires hospitalization/prolongation of hospitalization; disability=results in disability/incapacity; congen.=congenital anomaly/birth defect; other=other medically important information; (blank)=non-serious

 $^{2 \}quad \text{Drug role: } \textbf{S} \text{=-suspect; } \textbf{C} \text{=-concomitant; } \textbf{I} \text{=-interacting; } \textbf{N} \text{=-not administered}$

³ Additional Information on Drug: 1=Counterfeit; 2= Overdose; 3=Drug taken by the father; 4=Drug taken beyond expiry date; 5=Batch and lot tested and found within specifications; 6=Batch and lot tested and found not within specifications; 7=Medication error; 8=Misuse; 9=Abuse; 10=Occupational exposure; 11=Off label use; (blank) = no additional information