EVPM ICSR(s)	Individual Case Safety Report Form					EudraVigilance	
General Information							
EudraVigilance Local Report Number	EU-E	EU-EC-10012184028					
Sender Type	Regu	Regulatory authority					
Sender's Organisation	EEA I	EEA Regulator					
Type of Report	Spon	Spontaneous					
Primary source country	Europ	European Economic Area					
Reporter's qualification	Healt	Healthcare Professional					
Case serious?	Yes	Yes					
Patient							
Age Group	Age Group		Age Group (as per reporter)			Sex	
18-64 Years					Female		
Reaction / Event							
MedDRA LLT		Duration		Outcome		Seriousness ¹	
Debility			Red	Recovered/Resolved			
Vaccination site pain	·		Red	Recovered/Resolved			
Chest pain			Recovered/Resolved				
Missed abortion				Not Recovered/Not Resolved		other	
Vaccination site oedema				Recovered/Resolved			
Myalgia				Recovered/Resolved			
Fever		Recovered/Resolved					
Drug Information							
Role ² Drug	Drug		Dose		Units in Interval	Action taken	
S SPIKEVAX - ELASOMERAN							
Drug Information (cont.)							
Info ³ Drug	Drug		Indication		Pharm. Form	Route of Admin.	

N/A

SPIKEVAX - ELASOMERAN

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 Drug role: S=suspect; C=concomitant; I=interacting; N=not administered

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