General	Information

EudraVigilance Local Report Number EU-EC-10010893489

Sender Type Health professional

Sender's Organisation PFIZER S.R.L.

Type of Report Spontaneous

Primary source country Non-European Economic Area

Reporter's qualification Healthcare Professional

Case serious? Yes

Patient

. acronic		
Age Group	Age Group (as per reporter)	Sex
3-11 Years		Female

Reaction / Event						
MedDRA LLT		Duration	Outcome		Seriousness ¹	
Short of	breath		Recovered/Resolved			
Off label	use		Unknown			
Chest pa	Chest pain Recovered/Re		ed/Resolved			
Product use in unapproved population			Unknown			
Myopericarditis			Recovered/Resolved		other	
Drug Information						
Role ²	Drug	Duration	Dose	Units in Interval	Action taken	
S	TOZINAMERAN - TOZINAMERAN	1.0 Days		Total	Not applicable	

Drug I	Drug Information <i>(cont.)</i>			
Info ³	Drug	Indication	Pharm. Form	Route of Admin.
	TOZINAMERAN - TOZINAMERAN	COVID-19 immunisation		

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¹ 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