															CIC	)IVI	5 F	OF	RM
SUSPECT ADVERSE REACTION REPORT														—					_
SUSPE	SI ADVERSE F	REACTIO	N REPO	ΚI															
														Τ					
														$\perp$		L			
			I. REA	CTION	INFOR	MATION													
1. PATIENT INITIALS	1a. COUNTRY											8-12			K ALL		_		
(first, last) PRIVACY	DOMINICAN REPUBLIC	· I	vACY Year	7 Years	Male	Unk	Day		Month Unk		Year				OPRIA RSE R				
7 + 13 DESCRIBE REACTION(S) (including relevant tests/lab data) Event Verbatim [LOWER LEVEL TERM] (Related symptoms if any separated by commas)												PATIENT DIED							
problems with the device [Device defective]												INVOLVED OR							
Casa Description: This is a spontaneous report received from a Consumer or other non-LICE from a resident												PROLONGED INPATIENT HOSPITALISATION							
Case Description: This is a spontaneous report received from a Consumer or other non HCP from product quality group, Program ID: 164974.																			
A 7-year-old male patient received somatropin (GENOTROPIN PEN), (Batch/Lot number: unknown). The											INVOLVED PERSISTENT OR SIGNIFICANT DISABILITY OR								
patient's relevant medical history and concomitant medications were not reported.														ACITY					
The following information was reported: DEVICE DEFECTIVE (non-serious), outcome "unknown", described																			
as "problems with the device".  (Continued on Additional Information									ion F	age)	[	]	FE HRE	ATENII	NG				
		II.	SUSPEC	T DRU	IG(S) IN	FORMA	TION	1											
II. SUSPECT DRUG(S) INFORMATION  14. SUSPECT DRUG(S) (include generic name)												20. DID REACTION ABATE AFTER STOPPING							
#1 ) Genotropin Pen (SOMATROPIN) Solution for injection #2 ) Genotropin Pen (SOMATROPIN (DEVICE CONSTITUENT)) Solution for injection													DRUG		IEKS	IOPI	PING		
#2 ) Genotropin Pen (SOMATROPIN (DEVICE CONSTITUENT)) Solution for injection  15. DAILY DOSE(S)  16. ROUTE(S) OF ADMINISTRATION											┨	_		_	_	_			
#1) #2)						1 ) Unknown 2 ) Unknown							Y	ES	NO	) <b>[</b>	<b>N</b> A		
#2 ) #2 17. INDICATION(S) FOR USE						, Children							DID RI						
#1 ) Unknown #2 ) Unknown											R AFT DUCTI								
18. THERAPY DATES(fro	19. THERAPY	. THERAPY DURATION																	
#1 ) Unknown						1 ) Unknown							YES NO NA						
#2 ) Unknown	#2 ) Unkno	) Unknown								—									
		III. Co	DNCOMIT	ANT D	RUG(S	) AND H	ISTO	)R\	1										
22. CONCOMITANT DRU	JG(S) AND DATES OF ADM	IINISTRATION (	exclude those use	ed to treat re	eaction)														
														_		_			
23. OTHER RELEVANT I From/To Dates	HISTORY. (e.g. diagnostics,		ancy with last mo History / Notes	nth of perior	d, etc.) Description														
Unknown																			
		۱۱	. MANUF	ACTU	RER INI	FORMAT	ION												
24a. NAME AND ADDRESS OF MANUFACTURER Pfizer S.A.						MARKS													
Laura Arce Mora Avenida Escazú, T																			
San Jose, COST																			
	25b. NA	25b. NAME AND ADDRESS OF REPORTER								_		_			_				
	PV20250	0082181	NAME	NAME AND ADDRESS WITHHELD.															
24c. DATE RECEIVED BY MANUFACTURE	24d. REPORT	SOURCE	LITERATURE																
05-JUL-2025	LI STORY																		
	HEALTH		OTHER: Sponta	ai leous															
DATE OF THIS REPORT 21-JUL-2025	25a. REPORT		FOLLOWUP:																

## **ADDITIONAL INFORMATION**

## 7+13. DESCRIBE REACTION(S) continued

The action taken for somatropin was unknown.

Causality for "problems with the device" was determined associated to device constituent of somatropin (malfunction).

No follow-up attempts are possible. Batch/lot number is not provided, and it cannot be obtained.