													CIC	MS	FOI	RM	
SUSPECT ADVERSE REACTION REPORT																 	
		L DEA	OTION.	INFOR	NAATION		Ш								<u> </u>		
1. PATIENT INITIALS	1a. COUNTRY	I. KEAC	2a. AGE	3. SEX	MATION 3a. WEIGHT	_	6 DE	ACTION	ONSE	т	8-12	CHE	CK ALL				
(first, last) PRIVACY	DOMINICAN REPUBLIC	Day Month Year PRIVACY	14 Years	Male	Unk	Day		Month	Y	ear 025	APPROPRIATE TO						
7+13 DESCRIBE REACTION(S) (including relevant tests/lab data) Event Verbatim [LOWER LEVEL TERM] (Related symptoms if any separated by commas) the dosage wasn't showing properly [Device image display issue] previously it was 1.5 mg, but we were administering 1.4 mg because 1.5 wasn't available [Incorrect dose administered by device] At that moment, she wasn't administering it because she was planning to restart [Intentionally missed dose] previously it was 1.5 mg, but we were administering 1.4 mg because 1.5 wasn't available [Drug dose prescribing error] Case Description: This is a spontaneous report received from a Consumer or other non HCP, Program ID:											PATIENT DIED INVOLVED OR PROLONGED INPATIENT HOSPITALISATION INVOLVED PERSISTENT OR SIGNIFICANT DISABILITY OR INCAPACITY						
164974. (Continued on Additional Information Page								age)	LIFE THREATENING								
		II. SUSPEC	T DRU	G(S) IN	FORMA	TIO	N										
14. SUSPECT DRUG(S) (include generic name) #1) Genotropin Pen (SOMATROPIN) Solution for injection #2) Genotropin Pen (SOMATROPIN (DEVICE CONSTITUENT)) Solution for injection											20. DID REACTION ABATE AFTER STOPPING DRUG?						
#1) 1.4 mg (prescribed 1.5 mg)					: Route(s) of administration 1) Unknown 2) Unknown							YES NO NA					
17. INDICATION(S) FOR USE #1) Unknown #2) Unknown										21. DID REACTION REAPPEAR AFTER REINTRODUCTION?							
#1) Unknown / 2025 #					THERAPY DURATION) Unknown) Unknown							YES NO NA					
		III. CONCOMIT	ANT D	RUG(S) AND H	IST	OR'	Y									
	.,	IINISTRATION (exclude those use allergies, pregnancy with last mor Type of History / Notes	ed to treat re	action)													
		IV. MANUF	ACTUF			ION	1										
24a. NAME AND ADDRESS OF MANUFACTURER Pfizer S.A. Laura Arce Mora Avenida Escazú, Torre Lexus, piso 7. Escazú San Jose, COSTA RICA					MARKS												
24c. DATE RECEIVED BY MANUFACTURE	24b. MFR CC PV20250 24d. REPOR' STUDY		NAME	25b. NAME AND ADDRESS OF REPORTER NAME AND ADDRESS WITHHELD. NAME AND ADDRESS WITHHELD.													
13-JUN-2025	HEALTH		aneous														
18-JUN-2025	25a. REPOR	FOLLOWUP:															

ADDITIONAL INFORMATION

7+13. DESCRIBE REACTION(S) continued

A 14-year-old male patient received somatropin (GENOTROPIN PEN), (Batch/Lot number: unknown) till 2025 at 1.4 mg (prescribed 1.5 mg)). The patient's relevant medical history and concomitant medications were not reported. The following information was reported: INTENTIONAL DOSE OMISSION (non-serious) with onset 2025, described as "At that moment, she wasn't administering it because she was planning to restart"; INCORRECT DOSE ADMINISTERED BY DEVICE (non-serious), PRODUCT PRESCRIBING ERROR (non-serious) all with onset 2025 and all described as "previously it was 1.5 mg, but we were administering 1.4 mg because 1.5 wasn't available"; DEVICE INFORMATION OUTPUT ISSUE (non-serious) with onset 2025, described as "the dosage wasn't showing properly". The action taken for somatropin was temporarily withdrawn in 2025.

Additional Information: Caregiver stated: "I was calling because I needed an applicator. I didn't know if it was due to humidity or something else, but I couldn't see the dosage". When asked about the dosage used, she explained: "The doctor had changed it. I was going to pick it up again. She had increased the dose to 2 mg; previously it was 1.5 mg, but we were administering 1.4 mg because 1.5 wasn't available. Now, I wasn't sure if the humidity from refrigeration was affecting it, but the dosage wasn't showing properly. At that moment, she wasn't administering it because she was planning to restart. She had been giving it based on what she believed was correct, turning the applicator as the nurse had previously instructed. But since around Holy Week, the dosage hadn't been visible. As the treatment was nearing its end, she decided to wait and see if the doctor would prescribe it again. After their appointment last week, when the doctor did prescribe it again, she decided to call to see if the device could be replaced. She also mentioned that if there was a pharmacy where she could buy it, she would do so without any problem".

Causality for "the dosage wasn't showing properly", "previously it was 1.5 mg, but we were administering 1.4 mg because 1.5 wasn't available" and "at that moment, she wasn't administering it because she was planning to restart" was determined associated to device constituent of somatropin (malfunction).