													CIO)MS	FO	RM	
SUSPECT ADVERSE REACTION REPORT											 T	— П	 T		T		
		I DEA	CTION	INICOD	MATION												
1. PATIENT INITIALS	1a. COUNTRY	2. DATE OF BIRTH	2a. AGE	3. SEX	MATION 3a. WEIGHT		REAG	CTION (ONSET	r	8-12	CHEC	CK ALL				
(first, last) PRIVACY	DOMINICAN REPUBLIC	Day Month Year PRIVACY	11 Years	Male	Unk	Day		Month Jnk	Ye	ear	APPROPRIATE TO ADVERSE REACTION						
7 + 13 DESCRIBE REACTION(S) (including relevant tests/lab data) Event Verbatim [LOWER LEVEL TERM] (Related symptoms if any separated by commas) the needle keeps leaking liquid [Device leakage]										PATIENT DIED INVOLVED OR							
Case Description: This is a spontaneous report received from a Consumer or other non HCP from product quality group, Program ID: 164974.											HOSPITALISATION						
An 11-year-old male patient received somatropin (GENOTROPIN PEN), (Batch/Lot number: unknown) at 1 mg daily, Device Lot Number: LD7549, Device Expiration Date: Mar2027.											INVOLVED PERSISTENT OR SIGNIFICANT DISABILITY OR INCAPACITY						
(Continued on Additional Information Page)										ge)	LIFE THREATENING						
II. SUSPECT DRUG(S) INFORMATION																	
14. SUSPECT DRUG(S) (include generic name) #1) Genotropin Pen (SOMATROPIN) Solution for injection #2) Genotropin Pen (SOMATROPIN (DEVICE CONSTITUENT)) Solution for injection {Lot # LD7549}											20. DID REACTION ABATE AFTER STOPPING DRUG?						
15. DAILY DOSE(S) #1) 1 mg, daily #2)					. 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					THERAPY DURATION) Unknown) Unknown							YES NO NA					
		III. CONCOMIT	TANT D	RUG(S	AND H	ISTO	RY	,									
23. OTHER RELEVANT I	,	IINISTRATION (exclude those us		d, etc.)													
From/To Dates Unknown		Type of History / Notes		Description													
		IV. MANUF	ACTU	RER INF	ORMAT	ION											
24a. NAME AND ADDRESS OF MANUFACTURER Pfizer S.A. Laura Arce Mora Avenida Escazú, Torre Lexus, piso 7. Escazú San Jose, COSTA RICA					ARKS												
	24b. MFR CC PV20250	NTROL NO. 00072402			ME AND ADDR												
24c. DATE RECEIVED BY MANUFACTURE		NAME AND ADDRESS WITHHELD. NAME AND ADDRESS WITHHELD.															
DATE OF THIS REPORT 26-JUN-2025	T HEALTH PROFES 25a. REPORT			\dashv													

ADDITIONAL INFORMATION

7+13. DESCRIBE REACTION(S) continued

The patient's relevant medical history and concomitant medications were not reported.

The following information was reported: DEVICE LEAKAGE (non-serious), described as "the needle keeps leaking liquid".

Additional Information: Patient's caregiver stated: "I'm calling because my son's Genotropin device, when I insert the Genotropin and then remove the needle, the needle keeps leaking liquid, I think the device has something that is not working properly. And I would not want to take the risk of continuing to use such an expensive medication only for it to go to waste". As of 24Jun2024, the person in charge of the patient said: "This week they sent me a nurse because the device, when I gave my son the injection, I waited about 10 to 15 seconds for all the liquid to come out, and when I removed it, the device kept leaking liquid, little drops of liquid, which it did not do it before. The nurse told me to change the needle, I changed it, but the needle kept leaking".

Causality for "the needle keeps leaking liquid" was determined associated to device constituent of somatropin (malfunction).