													CIC	OMS	3 F	OR	M		
SUSPEC	T ADVERSE REA	ACTION REPO	RT																
									Τ					П			_		
											Ш			Ш			_		
1. PATIENT INITIALS	1a. COUNTRY	I. REAC	CTION I	NFORMATION  3. SEX 3a. WEIGHT	_	6 DE	ACTION	N ON	SET	I g.	12	CHE	CK ALL				_		
(first, last) PRIVACY	PANAMA Da		6	-emale Unk	Day	<del>-</del>	Month Unk	T	Yea	_	8-12 CHECK ALL 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) painful injection [Injection site pain] the medication could not be administered [Drug dose omission by device] injection process was difficult because the pen was not friendly [Device difficult to use]									PATIENT DIED  INVOLVED OR PROLONGED INPATIENT HOSPITALISATION										
the needles of the pen were damaged [Needle issue] the pen turned with difficulty/dosing button was jammed, it did not turned/it doesn't screw it on properly [Device mechanical jam] 80% of the medication was spilled. [Device leakage] the numbers were not visible [Device image display issue]									INVOLVED PERSISTENT OR SIGNIFICANT DISABILITY OR INCAPACITY										
(Continued on Additional Information Page								e)	LIFE THREATENING										
		II. SUSPEC	T DRUG	G(S) INFORMA	101T	<u></u>													
14. SUSPECT DRUG(S) (include generic name) #1 ) Genotropin Pen (SOMATROPIN) Solution for injection #2 ) Genotropin Pen (SOMATROPIN (DEVICE CONSTITUENT)) Solution for injection {Lot # A143}										20	20. DID REACTION ABATE AFTER STOPPING DRUG?								
15. DAILY DOSE(S) #1 ) 0.8 mg, daily #2 )				6. ROUTE(S) OF ADMINISTRATION 21 ) Unknown 22 ) Unknown							YES NO NA								
17. INDICATION(S) FOR USE #1 ) Unknown #2 ) Unknown								21	21. DID REACTION REAPPEAR AFTER REINTRODUCTION?										
18. THERAPY DATES(from #1 ) Unknown #2 ) Unknown	#1	. THERAPY DURATION 1 ) Unknown 2 ) Unknown							YES NO NA										
		III. CONCOMIT	ANT DF	RUG(S) AND H	ISTO	DR'	Y												
22. CONCOMITANT DRU	G(S) AND DATES OF ADMINIS																		
23. OTHER RELEVANT H From/To Dates Unknown	IISTORY. (e.g. diagnostics, aller	gies, pregnancy with last mo Type of History / Notes		etc.) Description															
		IV. MANUF	ACTUR	ER INFORMAT	TION	l													
24a. NAME AND ADDRES Pfizer S.A. Laura Arce Mora Avenida Escazú, To San jose, COSTA	orre Lexus, piso 7. Esca			26. REMARKS															
	24b. MFR CONTR PV20250004			25b. NAME AND ADDR					).								_		
24c. DATE RECEIVED BY MANUFACTURE	24d. REPORT SOI	JRCE		NAME AND ADD	RESS	S WI	ТНН	ELD	).										
15-APR-2025	HEALTH PROFESSION		aneous	NAME AND ADDRESS WITHHELD.															
DATE OF THIS REPORT 21-APR-2025	25a. REPORT TYF	FOLLOWUP:																	

## ADDITIONAL INFORMATION

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

Case Description: This is a spontaneous report received from a Consumer or other non HCP and a Nurse, Program ID: 164974.

A 6-year-old female patient received somatropin (GENOTROPIN PEN), (Batch/Lot number: unknown) at 0.8 mg daily, Device Lot Number: A143, Device Expiration Date: May2025. The patient's relevant medical history and concomitant medications were not reported.

The following information was reported: INJECTION SITE PAIN (non-serious), outcome "unknown", described as "painful injection"; DRUG DOSE OMISSION BY DEVICE (non-serious), outcome "unknown", described as "the medication could not be administered"; DEVICE DIFFICULT TO USE (non-serious), outcome "unknown", described as "injection process was difficult because the pen was not friendly"; NEEDLE ISSUE (non-serious), outcome "unknown", described as "the needles of the pen were damaged"; DEVICE MECHANICAL ISSUE (non-serious), outcome "unknown", described as "the pen turned with difficulty/dosing button was jammed, it did not turned/it doesn't screw it on properly"; DEVICE LEAKAGE (non-serious), outcome "unknown", described as "80% of the medication was spilled."; DEVICE INFORMATION OUTPUT ISSUE (non-serious), outcome "unknown", described as "the numbers were not visible". The action taken for somatropin was unknown.

Causality for "painful injection", "the medication could not be administered", "injection process was difficult because the pen was not friendly", "the needles of the pen were damaged", "the pen turned with difficulty/dosing button was jammed, it did not turned/it doesn't screw it on properly", "80% of the medication was spilled." and "the numbers were not visible" was determined associated to device constituent of somatropin (malfunction).

Additional information: Reporter stated the needles of the pen were damaged and the medication could not be administered because the pen turned with difficulty and the numbers were not visible, so it was not possible to set the dose. Reporter mentioned that she had two daughters and the injection process was difficult because the pen was not friendly and this led to a painful injection and she did not know if it was because of the pen or a bad administration technique. Reporter also stated that she set the medication and it did not screw correctly so 80% of the medication was spilled. Nurse stated the dosing button was jammed, it did not turned.