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										<u> </u>							<u> </u>		
I. REACTION INFORMATION 1. PATIENT INITIALS 1a. COUNTRY 2. DATE OF BIRTH 2a. AGE 3. SEX 3a. WEIGHT 4-6 REACTION ONSET 8-12 CHECK ALL																			
1. PATIENT INITIALS (first, last)						u. WEIGHT	Day 27	у	Month FEB	Т	Year 202		,	APPI ADVI	CK ALL ROPRI ERSE I ENT DI	ATE T REAC			
7 + 13 DESCRIBE REACTION(S) (including relevant tests/lab data) Event Verbatim [LOWER LEVEL TERM] (Related symptoms if any separated by commas) Indurated abscess in the left buttock / the surrounding area is hardened / Inflammatory changes / Slightly red color [Injection site abscess] her 'waist' hurts, the pain is on both sides and radiates down the lumbar area [Pelvic pain]								INVOLVED OR PROLONGED INPATIENT HOSPITALISATION INVOLVED PERSISTENT OR SIGNIFICANT DISABILITY OR INCAPACITY											
Case Description: This Study report from HONDURAS was received by Adium via Asofarma a tu lado Patient Support Program (reference number: HN-ADIUM-HN-0086-20250228) on 28-FEB-2025 from a Consumer/Other Non-Health Prof regarding a Child 8 Years old Female patient who experienced Indurated abscess in the left buttock / the surrounding area is hardened /										LIFE THREATENING CONGENITAL ANOMALY									
				(C	ontinu	ed on Ac	dition	al Inf	ormat	ion F	Page	, [] '	ОТН	ER				
II. SUSPECT DRUG(S) INFORMATION																			
14. SUSPECT DRUG(S) (include generic name) #1) Eligard® (Leuprolide acetate) Injection, 45 milligram {Lot # 15109CUY; Exp.Dt. MAR-2026} #2) Eligard® Unspecified Device (Leuprolide acetate) Injection {Lot # (Continued on Additional Information Page)									20. DID REACTION ABATE AFTER STOPPING DRUG?										
#1) 45 milligram, q 6 month #1					ROUTE(S) OF ADMINISTRATION) Subcutaneous use) Unknown							YES NO NA							
17. INDICATION(S) FOR USE #1) Precocious Puberty (Precocious puberty) #2) Precocious Puberty (Precocious puberty)								21. DID REACTION REAPPEAR AFTER REINTRODUCTION?											
#1) 19-FEB-2025 / Ongoing #1						THERAPY DURATION) Unknown) Unknown) Unknown													
		III. CON	COMITANT	DRUG	3(S) A	AND H	HST	OR	Y										
22. CONCOMITANT DRUG(S) AND DATES OF ADMINISTRATION (exclude those used to treat reaction) #1) OMEGA 3 [FISH OIL] (FISH OIL) ; Unknown #2) DIGLET (COLECALCIFEROL) ; Unknown																			
23. OTHER RELEVANT HISTORY. (e.g. diagnostics, allergies, pregnancy with last month of period, etc.) From/To Dates Type of History / Notes Description Unknown to Ongoing Current Condition Allergy Food allergy (Food allergy) Lactose, nuts																			
IV. MANUFACTURER INFORMATION																			
24a. NAME AND ADDRESS OF MANUFACTURER TOlmar, Inc. Laura Pethick, PharmD 701 Centre Ave Fort Collins, CO 80526 UNITED STATES					. REMAR														
	24b. MFR CONTROL NO. 25hN057313 25b. NAME AND ADDRESS OF REPORTER NAME AND ADDRESS WITHHELD.																		
24c. DATE RECEIVED BY MANUFACTURE 19-MAR-2025	24d. REPOR' STUDY HEALTH PROFES	LITE	ERATURE HER: Study																
DATE OF THIS REPORT 24-MAR-2025 Distribution 25a. Report Type																			

ADDITIONAL INFORMATION

7+13. DESCRIBE REACTION(S) continued

Inflammatory changes / Slightly red color (Injection site abscess), during Eligard (Leuprolide acetate) 45 milligram therapy for Precocious puberty. The needle component of the constituent device was not identified in the report. The report was sent to Tolmar on 03-MAR-2025.

The patient's medical history and current conditions included Precocious puberty, Food allergy.

Concomitant medications included OMEGA 3 and OTHER THERAPEUTIC PRODUCTS.

On 25-AUG-2023, the patient began receiving Eligard 45 milligram, q 6 month via Subcutaneous use for Precocious puberty (Lot numbers and Expiration dates not provided). On 19-FEB-2025, the patient received Eligard 45 milligram, q 6 month via Subcutaneous use (Lot number: 15109CUY; Expiration date: MAR-2016). On 27-FEB-2025, 6 fays after the most recent dose of Eligard, the patient experienced burning, redness and hardening at injection site. She was taken to a private hospital, an hemogram was attached and the indication for medication was given, which was applied at the time. The evaluation of the doctor concluded that it was an indurated abscess in the left buttock, but that the medication was applied in the left upper quadrant and it was not seen as an application error. On an unknown date, the person in charge of the program made an in-person follow-up visit to assess the affected area, the physical examination showed hardening in the left upper quadrant, a measurement was taken and it was observed that it was within the insertion area, there were inflammatory changes, slightly red color in the area, no temperature elevation was palpated, but the surrounding area was hardened. No fever was reported. She continued oral antibiotic therapy. Corrective treatment included ceftriaxone, cefalexin and acetaminophen.

Relevant test results included:

27-FEB-2025 : Granulocyte count: 53 (Ref range: Not provided)

27-FEB-2025 : Haematocrit: 39.5 % percent (Ref range: Not provided)

27-FEB-2025: Haemoglobin: 12.3 gram per decilitre (Ref range: Not provided)

27-FEB-2025 : Lymphocyte count: 35 (Ref range: Not provided)

27-FEB-2025: Mean cell haemoglobin: 26.7 picogram (Ref range: Not provided)

27-FEB-2025 : Mean cell haemoglobin concentration: 31.1 gram per decilitre (Ref range: Not provided)

27-FEB-2025: Mean cell volume: 85.7 (Ref range: Not provided)

27-FEB-2025: MID: 12 (Ref range: Not provided)

27-FEB-2025 : Red blood cell count: 4.61 cubic millimetre (Ref range: Not provided) 27-FEB-2025 : White blood cell count: 13,000 cubic millimetre (Ref range: Not provided)

27-FEB-2025 : Platelet count: 241,000 cubic millimetre (Ref range: Not provided)

On an unknown date: Body temperature: No fever (Ref range: Not provided)

Action taken with Eligard in response to the event was Dose Not Changed. De-challenge and re-challenge were Not applicable. The outcome of Injection site abscess was Unknown.

The reporter did not assess the seriousness or causality of the event in relationship to Eligard.

On 19-MAR-2025, follow-up information was received by Adium Patient support program (reference number: HN-ADIUM-HN-0086-20250228) from a Consumer/Other Non-Health Prof and sent to Tolmar on 20-MAR-2025. New information included: Added new event of her waist hurts, the pain was on both sides and radiates down the lumbar area (Pelvic pain) and clinical details of event Injection site abscess, updated event outcome of event Injection site abscess from Unknown to Recovered/Resolved.

On an unspecified date, at an unknown time after the most recent dose of Eligard, the patient experienced her waist hurts and that on Friday patient's mother would take her to the laboratory for a General Urine Test, as she hopes that it was not a urinary tract infection, as the pain was on both sides and radiates down the lumbar area. As per reported, there was no longer any sign of infection at the application site, area without redness, no hardening. Action taken with Eligard in response to the events was Dose Not Changed. De-challenge and re-challenge were Not applicable. The outcome of Injection site abscess was Recovered/Resolved. The outcome of Pelvic pain was Unknown.

The reporter did not assess the seriousness or causality of the events in relationship to Eligard.

13. Lab Data

#	Date	Test / Assessment / Notes	Results	Normal High / Low				
1		Body temperature						
		No fever						
2	27-FEB-2025	Granulocyte count						

ADDITIONAL INFORMATION

13. Lab Data #	Date	Test / Assessi	ment / Notes	Results	Normal High / Low				
3	27-FEB-2025	Haematoc	rit	39.5 % percent					
Ū			range not provided	00.0 /0 po.00					
4	27-FEB-2025	Haemoglo	bin	12.3 gram per decilitre					
		Reference	range not provided						
5	27-FEB-2025	Lymphocy	te count						
		35, Units a	and Reference range not	provided					
6	27-FEB-2025	Mean cell	haemoglobin	26.7 picogram					
		Reference range not provided							
7	27-FEB-2025	Mean cell concentrat	haemoglobin ion	31.1 gram per decilitre					
		Reference range not provided							
8	27-FEB-2025	Mean cell	volume						
85.7, Units and Reference range not provided									
9 27-FEB-2025 Platelet cou			unt	241,000 cubic millimetre					
		Reference range not provided							
10 27-FEB-2025 Red blood			cell count	4.61 cubic millimetre					
		Reference	range not provided						
11	27-FEB-2025	White bloo	d cell count	13,000 cubic millimetre					
Reference range not provided									
14-19. SUSPECT DRUG(S) continued									
14. SUSPECT DR	UG(S) (include generic name)		15. DAILY DOSE(S); 16. ROUTE(S) OF ADMIN	17. INDICATION(S) FOR USE	18. THERAPY DATES (from/to); 19. THERAPY DURATION				
, .) (Leuprolide acetate) Regimen #2) Injection,	45 milligram, q 6 month; Subcutaneous use	Precocious Puberty (Precocious puberty)	25-AUG-2023 / Ongoing; Unknown				
	#2) Eligard® Unspecified Device (Leuprolide UNK; Unknown acetate) Injection {Lot # 15109CUY}; Regimen (Precocious puberty) Unknown Unknown #1								