SUSPECT ADVERSE REACTION REPORT																			
SV-Tolmar-TLM-202	25-03410																		
				I. REAC	TION	INFORI	MATION												
1. PATIENT INITIALS 1a. COUNTRY 2. DATE OF BIRTH 2a. AG							E 3. SEX 4-6 REACTION ONSET								8-12	CHE			
(first, last) GAM EL Day			Month	- Y	ears	Male	Day Month Yea					Year			TO A	ROPR DVER	SE		
GAIVI	Cont	18	Jan	1955												KEA	CTION	ı	
7+13 DESCRIBE REA		ng relevant t	ests/lab data	a)				ļ	- 1							PATIE	ENT DI	ED	
1) Death (Death (10 (- 19/Apr/2025)	, .	(1001190	5))														THREA		NG
(- 19/Api/2025)	- Falai														Н	INVO	LVED (OR	
																HOSE	PITALIZ	ATIC	PATIENT ON
																PERS	JLTS IN SISTEN	CE O	ıR
										SIGNIFICANT DISABILITY/INCAPACITY									
										CONGENITAL ANOMALY OTHER MEDICALLY									
														LY NDITION					
				. SUSPECT	r dru	G(S)INF	ORMAT	ION											
14. SUSPECT DRUG(,	,													20.		EVEN		
1) Eligard® (Leuprolide acetate, Leuprolide acetate) (Suspect) (Injection)(Unk						nknown	1)						Cor	nt		ABA1 STOF	E AF	TER DRI	UG?
													COI	ιι		YES		NO	N
							ROUTE(S) OF ADMINISTRATION subcutaneous										EVENT PPEAF		
1) (45 milligram(s), 1 in 6 Month)						i) Subc	rculai ICUUS									AFTER REINTRODUCTION			
																YES		NO	N
17. INDICATION(S) FO	OR USE													_	(N	A : No	ot App	olica	ble)
1) prostate cancer [tate cance	r]																
18. THERAPY DATE(S) (from/to) 1) (07/Jun/2024 -) 19. THERAPY DURATION																			
				ONCOMITA	ANT D	DLIC(S)	V V VID HI	STOD)	,										
22. CONCOMITANT D	RUG(S) AND DAT	ES OF ADM				· ,			1										
No concomitants us	ed/reported																		
23. OTHER RELEVAN 1) PROSTATE CAN																			
			Г	V. MANUFA	ACTUF	RER INF	ORMAT	ION											
24a. NAME AND ADDRESS OF MANUFACTURER							Study Information												
Name : Tolmar, Inc 701 Centre Avenue							Study Name: NA												
Fort Collins, CO, 80526, UNITED STATES OF AMERICA							EudraCT Number: Protocol No.: NA												
Anjan.Chatterjee@tolmar.comand+1-9702124900							Center No.:												
							Sub	oject Id	:										
24.REPORT NULLIFIE	1	241	o. MFR CON	ITROL NO.															
L YES L	NO	sv	/-Tolmar-T	LM-2025-03	3410														
24c. DATE RECEIVED			d. REPORT																
BY MANUFACTU	IRER	I⊵	STUDY	LITE	RATURE	Ē													
13/Jun/2025				ROFESSIONAL															
DATE OF THIS REPO	RT		a. REPORT	TYPE															
17/Jun/2025			INITIAL	FOLI	LOWUP														

= Continuation attached sheet(s)..

Continuation Sheet for CIOMS report

1a. COUNTRY

EL SALVADOR

7+13 DESCRIBE REACTION(S) (including relevant tests/lab data) (Continuation...)

Event Description:

This study report from EL SALVADOR was received by Adium Pharma S.A through the Jazz Safety tool of the Patient Support Program "ASOFARMA A TU LA" (reference number: SV-ADIUM-SV-0007-20250613) on 13-Jun-2025, from a consumer (caregiver) (non-healthcare professional) regarding an elderly male patient who experienced serious event of "death" (Death) during Eligard (leuprolide acetate) 45 mg therapy for prostate cancer. The needle component of the constituent device was not identified in the report. The report was sent to Tolmar on 13-Jun-2025.

The patient's medical history was unknown and current condition included prostate cancer.

Concomitant medications were unknown.

On 07-Jun-2024, the patient began receiving Eligard 45 mg every 6 months via subcutaneous route for prostate cancer (Lot numbers and expiration dates were unknown).

On 19-Apr-2025, the patient died due to unknown cause of death. The patient was 70 years old at the time of his death. It was unknown if an autopsy was performed. No further details were provided.

Action taken with Eligard in response to event was not applicable. De-challenge and Re-challenge were not applicable.

The outcome of death was fatal.

The reporter assesses the seriousness of death as serious (death).

The reporter did not provide the causality of death in relationship to Eligard and Eligard Unspecified Device.

No further query was raised.

Listedness:

Death>Eligard>Unlisted as per CCDS>07-Nov-2024
Death>Eligard>Unlisted as per USPI>Feb-2025
Death>Eligard unspecified device>Unlisted as per USPI>Feb-2025
Death>Eligard>Unlisted as per Canadian monograph>02-Apr-2025

Company Remarks (Sender's Comments) :

Evaluator Comment (Tolmar): This is regarding 70-year-old elderly male patient who experienced fatal event of death (Death) while on Eligard(Leuprolide acetate) 45mg therapy for prostate cancer. Tolmar assessed death as serious due to fatal outcome. Causal role of Eligard (drug component) in the patient's death was not assessable as lack of information regarding cause of death, events or circumstances preceding patient's death, autopsy report, relevant medical history, supportive investigations that could indicate cause of death, concomitant medications received by the patient precludes meaningful medical assessment of the report. However, advanced age of the patient and underlying prostate cancer were pre-existing risk factors for the patient's death. Death was assessed as not related to the device component of Eligard.

14.SUSPECT DRUG(S) (Continuation...)

Product-Reaction Level

1) Drug : Eligard® (Leuprolide acetate)

Active Substance : 1) Leuprolide acetate

Drug Characterization : Suspect
Form of Admin : 1) Injection
Lot Number : 1) Unknown

Daily Dose : (45 milligram(s), 1 in 6 Month)

Route of Admin : 1) Subcutaneous

Indications : 1) prostate cancer [10060862 - Prostate cancer] Therapy Dates : 1) From : 07/Jun/2024 To :Not applicable

Action(s) Taken With Drug : Not applicable

Causality

1) Death (Death - 10011906, Death - 10011906)

Causality as per reporter : Not Related

Mfr. CONTROL NO: SV-Tolmar-TLM-2025-03410

Continuation Sheet for CIOMS report

Causality as per Mfr : Not assessable DeChallenge : Not applicable ReChallenge : Not Applicable

Labeling:

1) Death

CORE UnLabeled

2) Drug : Eligard® Unspecified Device (Leuprolide acetate)

Active Substance : 1) Leuprolide acetate

Drug Characterization : Suspect
Form of Admin : 1) Injection
Lot Number : 1) Unknown

Indications : 1) prostate cancer [10060862 - Prostate cancer]

Action(s) Taken With Drug : Not applicable

Causality

1) Death (Death - 10011906, Death - 10011906)
Causality as per reporter : Not Related
Causality as per Mfr : Not Related
DeChallenge : Not applicable
ReChallenge : Not Applicable

Labeling :

1) Death
CORE