SUS	PECT ADVERSI	E REACTION	ON REPOR	RT																
PA-Tolmar-TLM-202	25-04091																			
				I REAC	CTION	INFORM	MATION													
1. PATIENT INITIALS 1a. COUNTRY 2. DATE OF BIRTH 2a. AGE														8-12 CHECK ALL						
(first, last) PANAMA Day Month Year 75							Male	Day	ay Month		Year] 	TO A	ROPR DVEF	RSE			
L-J	I AINAWA	29	Sep	1949		75	Iviaic	24		Jun	۱	2	2025	REACTION						
7+13 DESCRIBE REA	. , .	•		•												PATIE	ENT DI	ED		
1) Increased Prostatic Antigen values (Prostatic specific antigen increased (10036975), Prostatic specific antigen increased (10036975))									reas	ed		LIFE	THREA	ATEN	ING					
(24/Jun/2025 -)	- Not Recovered	/Not Resol	ved/Ongoi	ng											느	INVO	LVED	OR		
Cont							nt	PROLONGED INPATIENT HOSPITALIZATION					Τ							
									RESULTS IN PERSISTENCE OR SIGNIFICANT											
									DISABILITY/INCAPACITY											
									CONGENITAL ANOMALY					1						
										R MEI		LLY NDITION	N							
			II	. SUSPEC	T DRU	G(S)INF	ORMAT	ION												
14. SUSPECT DRUG(,	,				. ,									20.	DID E				_
1) Eligard® (Leuprol	lide acetate, Leu	prolide ace	etate) (Sus _l	pect) (45 M	lilligran	n, Injecti	ion)(Unkr	nown)					Cor	nt		ABA1 STOR	E AF	TER DR	UG?	
																YES		NO	Δ	NA
15. DAILY DOSE(S)						16. ROUTE(S) OF ADMINISTRATION]:	21.	DID E						
1) (45 milligram(s), 1	1 in 6 Month)					1) Subc	Subcutaneous REAPPEAR AFTER AFTER REINTRODUC								OUC	ΓΙΟΝ				
																YES		NO	\square	NA
17. INDICATION(S) FO	AD LIGE													႕	(N	IA : No	ot Ap	olica	ble)	
1) Prostate cancer [tate cance	r]																	
18. THERAPY DATE(S (26-Jan-2024 - Ongo	THERAPY DATE(S) (from/to) 19. THERAPY DURATION 3- Jan-2024 - Ongoing)																			
(9)			ONOONIT	ANTO	DU (0)		2700	,											_
22. CONCOMITANT D	RUG(S) AND DAT	ES OF ADM		ONCOMITA ON (exclude t		. ,			<u>′</u>											
No concomitants us		20 01 71211		on (exclude)		oca to tre	at rouotion	''												
23. OTHER RELEVAN 1) PROSTATE CAN						nth of pe	eriod, etc.)													
I) I ROSTATE CAN	ICEN (10000002	, i iostate	cancer) (C	ontinuing.	163)															
				√. MANUF	ACTUF	RER INF	ORMATI	ON												
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	ject Id	:											
24.REPORT NULLIFIED 24b. MFR CONTROL NO.																				
L YES L	NO	PA	-Tolmar-Ti	_M-2025-0 ²	4091															
24c. DATE RECEIVED)		. REPORT		.001															
BY MANUFACTU	RER		STUDY	LITE	RATURE	Ē														
24/Jun/2025 HEALTH PROFESSIONAL																				
DATE OF THIS REPO	RT	l	a. REPORT	TYPE																
27/Jun/2025 INITIAL FOLLOWUP																				

= Continuation attached sheet(s)..

Mfr. CONTROL NO :PA-Tolmar-TLM-2025-04091

Continuation Sheet for CIOMS report

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

Event Description:

This study report from Panama was received by Adium (reference number: PA-ADIUM-PA-0068-20250624) via Patient Support Program on 24-Jun-2025 from a consumer (patient or family member) (non-healthcare professional) regarding an elderly 75-year-old male patient who experienced non-serious event of "Increased Prostatic Antigen values" (prostatic specific antigen increased) 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 25-Jun-2025.

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

Concomitant medication was unknown

On 26-Jan-2024, the patient began receiving Eligard 45 mg, every 6 months via subcutaneous route for prostate cancer (Lot number and Expiration dates were not provided).

On 24-Jun-2025, the patient's prostate-specific antigen values increased severely with a result of 76.

Corrective treatment included to start bicalutamide as doctor instructed.

Relevant test results included:

On 24-Jun-2025: PSA: increased prostatic antigen with a result of 76 (Ref range: not provided).

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

The outcome of prostatic specific antigen increased was not recovered.

The reporter did not assess the seriousness of prostatic specific antigen increased.

The reporter assessed the causality of prostatic specific antigen increased in relationship to Eligard and Eligard unspecified device as related.

No further information is expected as the reporter did not consent to be contacted for follow-up.

Listedness:

Prostatic specific antigen increased>Eligard>listed as per CCDS>07-Nov-2024 Prostatic specific antigen increased>Eligard>listed as per USPI>Feb-2025

Prostatic specific antigen increased>Eligard unspecified device>listed as per USPI>Feb-2025

Prostatic specific antigen increased>Eligard>listed as per Canadian monograph>02-Apr-2

Company Remarks (Sender's Comments):

Evaluator comment (Tolmar): This is regarding an elderly 75-year-old female patient who had prostate specific antigen increased (Increased Prostatic Antigen values), during Eligard (leuprolide acetate) 45 mg therapy for prostate cancer. Tolmar assessed the event as non-serious since it does not meet the ICH seriousness criteria and is not an IME event. The reported event was assessed as not related to Eligard (drug and device) as the event can be explained by underlying prostate cancer.

Additional Information (Continuation...)

Lab Result:

Test Name	Test Date	Test Result	Normal Value
PSA	24/Jun/2025		

Test Result (Code) / Result Unstructured Data (free text) :

1) Test Name: PSA

Result Unstructured Data (free text): Increased Prostatic Antigen with a result of 76

Test Date: 24/Jun/2025

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

Product-Reaction Level

1) Drug : Eligard® (Leuprolide acetate)
Active Substance : 1) Leuprolide acetate

Continuation Sheet for CIOMS report

Drug Characterization : Suspect
Form Strength : 1) 45 Milligram
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]

Action(s) Taken With Drug : Dose not changed

Causality

1) Increased Prostatic Antigen values (Prostatic specific antigen increased - 10036975, Prostatic specific antigen increased - 10036975)

Causality as per reporter : Related
Causality as per Mfr : Not Related
DeChallenge : Not applicable
ReChallenge : Not Applicable

Labeling:

1) Increased Prostatic Antigen values

CORE Labeled

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

Active Substance : 1) Leuprolide acetate

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

2) Unknown

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

Action(s) Taken With Drug : Not applicable

Causality

1) Increased Prostatic Antigen values (Prostatic specific antigen increased - 10036975, Prostatic specific antigen increased - 10036975)

Causality as per reporter : Related
Causality as per Mfr : Not Related
DeChallenge : Not applicable
ReChallenge : Not Applicable

Labeling:

1) Increased Prostatic Antigen values

CORE

15. DAILY DOSE(S) (Continuation...)

Dosage Text : Drug 1 :Eligard®

1) 45 Milligrams every 6 Months