			o., 55505																
SUS	PECT ADVERSI	E REACTI	ON REPOR	RT															
CR-Tolmar-TLM-20	25-01849																		
				I DEAC	TION	INFOR	MATION	-				<u> </u>						<u> </u>	•
I. REACTION							SEX 4-6 REACTION ONSET					1	8-12 CHECK ALL						
(first, last) COSTA RICA			y Month Year			ears 93	Male	Day Month Yea			′ear	\dashv		APPR TO AL	OVERS	ATE SE			
VSA COSTARICA 04			Dec 1931			93	I wais					2022			REACTION				
7+13 DESCRIBE REA	ACTION(S) (includin	ng relevant	ests/lab data	1)				ļ						\dashv_{l}	П	PATIE	NT DIE	D	
1) diagnosed with p (Progression of pro				, ,		conditio	on has wo	rsened	/ de	cay in	his	conc	lition	ٔ ا		LIFE T	HREAT	ENIN	IG
Recovering/Res	•	300409), 1	TOSIAIC CAI	10000	3002))										ᆷ	I INVOL'	VED O	R	
2) Bone metastasis (Metastases to bone (10027452), Metastases to bone (10027 (//2022 -) - Not Recovered/Not Resolved/Ongoing					(10027	452))	52))							PROLONGED INPATIENT HOSPITALIZATION RESULTS IN					
3) Eligard was no longer available in the country (Product supply issue (10077801),), Produc	Product supply issue (10077801))							PERSISTENCE OR SIGNIFICANT					
Unknown 4) Therefore, patient was switched to Zalodex 10.80 (Product substitution (1007675)						753). Proc), Product substitution (10076753))							DISABILITY/INCAPACITY CONGENITAL ANOMALY					
Unknown								,			,,				R MEDI				
Cont.							nt	$\underline{\mathbf{M}}$	IMPOR	RTANT	CON	DITION							
			II.	. SUSPECT	T DRU	G(S)IN	FORMAT	ION											
14. SUSPECT DRUG(, ,	,	-1-1-) (0 -	() (1	C>/L		- \							20	:0.	DID E		FR	
Eligard® (Leuprolide acetate, Leuprolide acetate) (Suspect) (Injection)(Unkr					ınknowi	n)						Con	nt	\vdash	ABATE STOP				
15. DAILY DOSE(S) 16. ROU					JTE(S) OF	ADMINI	STR	ATION	1				 :1.	■YES DID E		10	NA		
1) (45 milligram(s),	1 in 6 Month)						I) Subcutaneous									REAP!	PEAR R		
					REINTRODUCTION														
															(N.	JYES A∶Not		≀o Iicab	NA ole)
17. INDICATION(S) FO		tate cance	arl												`				,
Prostate cancer [10060862 - Prostate cancer] Herapy DATE(S) (from/to) 19. THERAPY DURATION													\dashv						
(25-Oct-2024 -)																			
			III. C	ONCOMITA	ANT D	RUG(S) AND HI	STORY											
22. CONCOMITANT D		ES OF ADN	INISTRATIC	N (exclude f	those u	sed to tre	eat reaction	٦)											
No concomitants us	ed/reported																		
23. OTHER RELEVAN	IT HISTORY (e.g. c	liagnostics,	allergies, pre	gnancy with	last mo	onth of p	eriod, etc.)												
1) PROSTATE CAN	ICER (10060862	, Prostate	cancer) (Co	ontinuing: \	Yes)														
														—	—		—		
	DE00 05 MANUE			V. MANUFA	ACTUF	RER INI													
24a. NAME AND ADDRESS OF MANUFACTURER Name : Tolmar, Inc							Study Information Study Name: NA												
701 Centre Avenue Fort Collins, CO, 80526, UNITED STATES OF AMERICA						EudraCT Number:													
debbie.maierhofer@tolmar.comand+1-4129158447						Protocol No.: NA Center No.:													
							ı	nter No. Dject Id											
24.REPORT NULLIFIE	D	24	b. MFR CON	TROL NO.				•											
YES	NO) Tolmor Ti	M 2025 0	1040														
24c. DATE RECEIVED)		R-Tolmar-TL d. REPORT S		1849														
BY MANUFACTU	IRER		STUDY	Пите	RATURE	=													
21/May/2025				OFESSIONAL															
DATE OF THIS REPO	RT	25	a. REPORT 1	_															
28/May/2025			INITIAL	FOL	LOWUP														

= Continuation attached sheet(s)..

Continuation Sheet for CIOMS report

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

5) Eligard was no longer having the expected effect/ lack of efficacy (Lack of drug effect (10023610), Drug ineffective (10013709)(27/Apr/2025 -) - Unknown)

Event Description:

This study report from Brazil was received by Adium Patient Support Program (reference number: CR-ADIUM-CR-0166-20250514) on 14-May-2025 from a consumer (non-healthcare professional) regarding an elderly 93-year old male patient who experienced serious (medically significant) events of "Bone metastasis" (Metastases to bone) and "diagnosed with prostate cancer with bone metastasis/ my grandfather's condition has worsened/ decay in his condition" (Prostate cancer) and non-serious events of "Eligard was no longer having the expected effect" (Drug ineffective), "Eligard was no longer having the expected effect and was no longer available in the country" (Product supply issue), and "Therefore, patient was switched to Zalodex 10.8" (Product substitution) during Eligard (Leuprolide acetate) 45mg therapy for prostate cancer. The needle component of the constituent device was not identified in the report. The report was sent to Tolmar on 15-May-2025.

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

Concomitant medications were not provided.

On an unknown date, the patient began receiving Eligard of 45 mg strength in 6 months of frequency via subcutaneous route of administration for prostate cancer (Lot numbers and Expiration dates were not provided).

On an unknown date in 2022, the patient had been diagnosed with prostate cancer with bone metastasis for 3 years. He was unaware of his condition, so he was only told that the medication being administered is to keep his prostate healthy and allow him to urinate. In recent months, his condition had worsened, but he was now recovering. According to the latest tests performed, Eligard was no longer having the expected effect and was no longer available in the country.

On 25-Apr-2025, the patient was switched to Zalodex - 10.80 milligram dose - for prostate cancer.

Corrective treatment was not provided.

Action taken with Eligard in response to the events was unknown. De-challenge and re-challenge were not applicable.

The outcome of metastases to bone was not recovered, prostate cancer was recovering and drug ineffective, product substitution and product supply issue was unknown.

The reporter did not assess the seriousness of metastases to bone, prostate cancer, drug ineffective and product substitution and product supply issue.

The reporter assessed the causality of drug ineffective and prostate cancer as related, metastases to bone as not related, product substitution and product supply issue as not reported in relationship to Eligard and Eligard unspecified device

No further queries were raised.

On 19-May-2025, Tolmar case ID QE-027981 was added to the case.

On 21-May-2025, the follow up was received by Adium Patient Support Program "ASOFARMA A TU LADO" (reference number: CR-ADIUM-CR-0166-20250514). New information included: dose administration date and lab test were added. Narrative was updated.

On 25-Oct-2024, the patient began receiving Eligard of 45 mg strength in 6 months of frequency via subcutaneous route of administration for prostate cancer (Lot numbers and Expiration dates were not provided).

Relevant lab test

On 25-Apr-2025, Blood testosterone: Unknown (Ref. range not provided).

No further queries were raised.

On 23-May-2025, Tolmar Quality Assurance number QE-027981 was added. This follow up information which was received on 23-May-2025 was considered as non-significant case.

Listedness

Metastases to bone>Eligard® >listed as per CCDS Eligard®> 7-Nov-2024

Metastases to bone> Eligard® >listed as per Canadian Monograph Eligard®> 2-Apr-2025

Metastases to bone> Eligard®>listed as per USPI Eligard®>Feb-2025

Metastases to bone> Eligard® Unspecified Device> listed as per USPI Eligard®>Feb-2025

Prostate cancer>Eligard® >listed as per CCDS Eligard®> 7-Nov-2024
Prostate cancer> Eligard® >listed as per Canadian Monograph Eligard®> 2-Apr-2025

Mfr. CONTROL NO: CR-Tolmar-TLM-2025-01849

Continuation Sheet for CIOMS report

Prostate cancer> Eligard®>listed as per USPI Eligard®>Feb-2025

Prostate cancer> Eligard® Unspecified Device> listed as per USPI Eligard®>Feb-2025

Product supply issue >Eligard® >unlisted as per CCDS Eligard®> 7-Nov-2024

Product supply issue > Eligard® > unlisted as per Canadian Monograph Eligard® > 2-Apr-2025

Product supply issue > Eligard®>unlisted as per USPI Eligard®>Feb-2025

Product supply issue > Eligard® Unspecified Device>unlisted as per USPI Eligard®>Feb-2025

Product substitution >Eligard® >unlisted as per CCDS Eligard® > 7-Nov-2024

Product substitution> Eligard® >unlisted as per Canadian Monograph Eligard®> 2-Apr-2025

Product substitution> Eligard®>unlisted as per USPI Eligard®>Feb-2025

Product substitution> Eligard® Unspecified Device>unlisted as per USPI Eligard®>Feb-2025

Drug ineffective >Eligard® >listed as per CCDS Eligard®> 7-Nov-2024

Drug ineffective> Eligard® >listed as per Canadian Monograph Eligard®> 2-Apr-2025

Drug ineffective> Eligard®>listed as per USPI Eligard®>Feb-2025

Drug ineffective> Eligard® Unspecified Device>listed as per USPI Eligard®>Feb-2025

Listedness of the events metastases to bone, prostate cancer, product supply issue, product substitution and drug ineffective is retained as per previous assessment.

Company Remarks (Sender's Comments) :

Evaluator comment (Tolmar): The case is regarding an elderly 93-year old male patient who reported events of Metastases to bone (Bone metastasis), Prostate cancer (diagnosed with prostate cancer with bone metastasis/ my grandfather's condition has worsened/ decay in his condition), Drug ineffective (Eligard was no longer having the expected effect), Product supply issue (Eligard was no longer having the expected effect and was no longer available in the country), Product substitution (Therefore, patient was switched to Zalodex 10.8) during Eligard (Leuprolide acetate) 45mg therapy for prostate cancer. Tolmar assessed the events bone metastasis and prostate cancer as serious (medically significant) as these events are included in IME list and rest of the events were assessed as non-serious since they did not meet the ICH seriousness criteria. The causality of the events bone metastasis and prostate cancer were assessed as not related to suspect Eligard(Drug and device) as these events could be attributed to underlying prostate cancer which is known to progress despite treatment. The causality of event drug ineffective was assessed as related to suspect drug Eligard (not related to device) based on timeline association with Eligard administration and given that the effect of drug may vary from person to person. The causality of events product substitution due to product supply issue was assessed as not related to suspect Eligard(drug and device) as it was human action due circumstances.

FU-Causality of the events metastases to bone, prostate cancer, product supply issue, product substitution and drug ineffective is retained as per previous assessment.

Additional Information (Continuation...)

Lab Result :

Test Name	Test Date	Test Result	Normal Value
TESTOSTERONE	25/Apr/2025		

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

1) Test Name: TESTOSTERONE

Result Unstructured Data (free text): Unknown

Test Date: 25/Apr/2025

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]

Action(s) Taken With Drug : Unknown

Continuation Sheet for CIOMS report

Causality

1) diagnosed with prostate cancer with bone metastasis/ my grandfather's condition has worsened/ decay in his condition (Progression of prostate cancer - 10066489, Prostate cancer - 10060862)

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

2) Bone metastasis (Metastases to bone - 10027452, Metastases to bone - 10027452)

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

3) Eligard was no longer available in the country (Product supply issue - 10077801, Product supply issue - 10077801)

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

4) Therefore, patient was switched to Zalodex 10.80 (Product substitution - 10076753, Product substitution - 10076753)

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

5) Eligard was no longer having the expected effect/ lack of efficacy (Lack of drug effect - 10023610, Drug ineffective - 10013709)

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

Labeling:

1) diagnosed with prostate cancer with bone metastasis/ my grandfather's condition has worsened/ decay in his condition

Labeled CORE

2) Bone metastasis

CORE Labeled

3) Eligard was no longer available in the country CORF

UnLabeled

4) Therefore, patient was switched to Zalodex 10.80 CORF Unl abeled

5) Eligard was no longer having the expected effect/ lack of efficacy 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

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

Action(s) Taken With Drug Not applicable

Causality

1) diagnosed with prostate cancer with bone metastasis/ my grandfather's condition has worsened/ decay in his condition (Progression of prostate cancer - 10066489, Prostate cancer - 10060862)

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

2) Bone metastasis (Metastases to bone - 10027452, Metastases to bone - 10027452)

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

3) Eligard was no longer available in the country (Product supply issue - 10077801, Product supply issue - 10077801)

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

4) Therefore, patient was switched to Zalodex 10.80 (Product substitution - 10076753, Product substitution - 10076753)

Continuation Sheet for CIOMS report

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

5) Eligard was no longer having the expected effect/ lack of efficacy (Lack of drug effect - 10023610, Drug ineffective - 10013709)

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

Labeling:

- 1) diagnosed with prostate cancer with bone metastasis/ my grandfather's condition has worsened/ decay in his condition CORE
- 2) Bone metastasis

CORE

3) Eligard was no longer available in the country CORE

4) Therefore, patient was switched to Zalodex 10.80

CORE

5) Eligard was no longer having the expected effect/ lack of efficacy

CORE