SUSPECT ADVERSE REACTION REPORT																				
DO-Tolmar-TLM-20	25-01817																			
				I. REAC	TION	INFORI	MATION													
1. PATIENT INITIALS		3. SEX 4-6 REACTION ONSET							8-12	CHE										
(first, last)	DOMINICAN Day Month Year						Male	Day Month Year					⁄ear			TO A	ROPF	RSE	E	
FRE	Oct	1949	75				2025			2025			KEA	CTIO	IN					
7+13 DESCRIBE REACTION(S) (including relevant tests/lab data)  1) Rash/rash on his upper limbs (Localised rash (10062704), Rash (10037844)) (//2025 - ) - Recovered/Resolved  2) Bilateral breast swelling (Breast swelling (10006312), Breast swelling (10006312)) (//2025 - ) - Recovered/Resolved  II. SUSPECT DRUG(S)INFORMATION														PATIENT DIED  LIFE THREATENING  INVOLVED OR PROLONGED INPATIENT HOSPITALIZATION RESULTS IN PERSISTENCE OR SIGNIFICANT DISABILITY/INCAPACITY CONGENITAL ANOMALY OTHER MEDICALLY IMPORTANT CONDITION  20. DID EVENT						
1) Eligard® (Leuprolide acetate, Leuprolide acetate) (Suspect) (22.5 Milligram, Injection)(15109CUY;UNK;UNK)  Cont.											nt		ABAT STOI YES	PPIN	TEF G DF NO	RUG	? NA			
							S. ROUTE(S) OF ADMINISTRATION									DID E				
1) (22.5 milligram(s), 1 in 3 Month)						1) Subo	ocutaneous								[ (N	AFTE REIN YES	ER ITRO	DUC No	Ş	NA
17. INDICATION(S) FOR USE 1) Prostate cancer [10060862 - Prostate cancer]															(		0171	γριιο	abio	,
1	8. THERAPY DATE(S) (from/to)  -Feb-2025 - Ongoing)  19. THERAPY DURATION																			
			III C	ONCOMITA	ANT D	RUG(S	) AND HIS	STORY	,											
22. CONCOMITANT D No concomitants us 23. OTHER RELEVAN	ed/reported		IINISTRATIO	ON (exclude t	hose us	sed to tre	eat reaction													
1) PROSTATE CAN																				
			ľ	V. MANUFA	ACTUF	RER INF	FORMATI	ION												
24a. NAME AND ADDRESS OF MANUFACTURER Name: Tolmar, Inc 701 Centre Avenue Fort Collins, CO, 80526, UNITED STATES OF AMERICA							Study Information Study Name: NA EudraCT Number: Protocol No.: NA Center No.: Subject Id:													
24.REPORT NULLIFIE  YES  24c. DATE RECEIVED	NO	DC	o. MFR CON O-Tolmar-T	LM-2025-0	1817															
BY MANUFACTU			STUDY		RATURE	Ē														
13/May/2025 HEALTH PROFESSIONAL																				
DATE OF THIS REPO 22/May/2025																				
ZZIIVIAYIZUZU			INITIAL	FOLI	LOWUP															

= Continuation attached sheet(s)..

### Continuation Sheet for CIOMS report

## 1a. COUNTRY

#### DOMINICAN REPUBLIC

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

#### Event Description :

This study report from Dominican Republic was received by Adium via Patient Support Program (reference number: DO-ADIUM-DO-0037-20250513) on 13-May-2025 from a consumer (non-healthcare professional) regarding an elderly 75-year-old male patient who experienced non serious events of "rash/rash on his upper limbs" (rash) and "bilateral breast swelling" (breast swelling) during Eligard (leuprolide) 22.5 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 14-May-2025.

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

Concomitant medications were unknown.

On an unknown date in Feb-2025, the patient began receiving Eligard 22.5 mg, every 3 months via subcutaneous route for prostate cancer (Lot numbers; 15109CUY; UNK; UNK and Expiration dates; May-2026; UNK; UNK).

On an unknown date in 2025, the patient experienced rash on his upper limbs which was moderate in nature. Additionally, he also had mild bilateral breast swelling which disappeared without further action.

Corrective treatment included topical hydrocortisone cream for rash.

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

The outcome of rash and breast swelling was recovered.

The reporter did not assess the seriousness or causality of rash, breast swelling in relationship to Eligard and Eligard unspecified device.

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

#### Listedness:

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

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

## Company Remarks (Sender's Comments):

Evaluator comment (Tolmar): This is regarding an elderly 75-year-old male patient who developed rash and breast swelling during Eligard (leuprolide acetate) 22.5 mg therapy for prostate cancer. Tolmar assessed the reported events as non-serious since they did not meet the ICH seriousness criteria and are not IME events. The reported events were assessed as related to Eligard (drug) based on plausible temporality and as not related to device.

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

#### Product-Reaction Level

1) Drug : Eligard® (Leuprolide acetate)

Active Substance : 1) Leuprolide acetate

Drug Characterization : Suspect
Form Strength : 1) 22.5 Milligram

Form of Admin : 1) Injection

Lot Number : 1) 15109CUY;UNK;UNK
Daily Dose : (22.5 milligram(s), 1 in 3 Month)

Route of Admin : 1) Subcutaneous

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

Action(s) Taken With Drug : Dose not changed

## Causality

# Continuation Sheet for CIOMS report

1) Rash/rash on his upper limbs (Localised rash - 10062704, Rash - 10037844)

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

2) Bilateral breast swelling (Breast swelling - 10006312, Breast swelling - 10006312)

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

Labeling:

1) Rash/rash on his upper limbs

CORE UnLabeled

2) Bilateral breast swelling

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) 15109CUY;UNK;UNK

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

Action(s) Taken With Drug : Not applicable

Causality

1) Rash/rash on his upper limbs (Localised rash - 10062704, Rash - 10037844)

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

2) Bilateral breast swelling (Breast swelling - 10006312, Breast swelling - 10006312)

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

Labeling:

1) Rash/rash on his upper limbs

CORE

2) Bilateral breast swelling

CORE

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

Dosage Text : Drug 1 :Eligard®

1) Expiration date: May-26;UNK;UNK ELIGARD 22.5 MG x 1 LIO x 2 JER

Drug 2 :Eligard® Unspecified Device
1) Expiration date: May-26;UNK;UNK