

<b>SUSPECT ADVERSE REACTION REPORT</b>  SV-Tolmar-TLM-2025-04833												

## I. REACTION INFORMATION

1. PATIENT INITIALS (first, last)	1a. COUNTRY	2. DATE OF BIRTH			2a. AGE Years	3. SEX	4-6 REACTION ONSET			8-12 CHECK ALL APPROPRIATE TO ADVERSE REACTION
NPP	EL	Day 26	Month Nov	Year 1950		Male	Day	Month	Year	
Cont..										
7+13 DESCRIBE REACTION(S) (including relevant tests/lab data) 1) Death (Death (10011906), Death (10011906)) ( - //2025) - Fatal										<input checked="" type="checkbox"/> PATIENT DIED <input checked="" type="checkbox"/> LIFE THREATENING <input checked="" type="checkbox"/> INVOLVED OR PROLONGED INPATIENT HOSPITALIZATION <input checked="" type="checkbox"/> RESULTS IN PERSISTENCE OR SIGNIFICANT DISABILITY/INCAPACITY <input type="checkbox"/> CONGENITAL ANOMALY <input type="checkbox"/> OTHER MEDICALLY IMPORTANT CONDITION

## II. SUSPECT DRUG(S) INFORMATION

14. SUSPECT DRUG(S)(include generic name) 1) Eligard® (Leuprolide acetate, Leuprolide acetate) (Suspect) (Injection)(Unknown)		20. DID EVENT ABATE AFTER STOPPING DRUG? <input type="checkbox"/> YES <input type="checkbox"/> NO <input checked="" type="checkbox"/> NA  21. DID EVENT REAPPEAR AFTER REINTRODUCTION <input type="checkbox"/> YES <input type="checkbox"/> NO <input checked="" type="checkbox"/> NA (NA : Not Applicable)	
15. DAILY DOSE(S) (45 milligram(s))			
16. ROUTE(S) OF ADMINISTRATION 1) Subcutaneous			
17. INDICATION(S) FOR USE 1) prostate cancer [10060862 - Prostate cancer]			
18. THERAPY DATE(S) (from/to) 1) (27/Nov/2021 - )		19. THERAPY DURATION	

## III. CONCOMITANT DRUG(S) AND HISTORY

22. CONCOMITANT DRUG(S) AND DATES OF ADMINISTRATION (exclude those used to treat reaction) No concomitants used/reported
23. OTHER RELEVANT HISTORY (e.g. diagnostics, allergies, pregnancy with last month of period, etc.) 1) PROSTATE CANCER (10060862, Prostate cancer) (27/Nov/2021 - ) (Continuing: No)

## IV. MANUFACTURER INFORMATION

24a. NAME AND ADDRESS OF MANUFACTURER Name : Tolmar, Inc 701 Centre Avenue Fort Collins, CO, 80526, UNITED STATES OF AMERICA Anjan.Chatterjee@tolmar.comand+1--9702124900		Study Information Study Name: NA EudraCT Number: Protocol No.: NA Center No.: Subject Id :	
24. REPORT NULLIFIED <input type="checkbox"/> YES <input type="checkbox"/> NO	24b. MFR CONTROL NO. SV-Tolmar-TLM-2025-04833		
24c. DATE RECEIVED BY MANUFACTURER 18/Jul/2025	24d. REPORT SOURCE <input checked="" type="checkbox"/> STUDY <input type="checkbox"/> LITERATURE <input type="checkbox"/> HEALTH PROFESSIONAL		
DATE OF THIS REPORT 24/Jul/2025	25a. REPORT TYPE <input checked="" type="checkbox"/> INITIAL <input type="checkbox"/> FOLLOWUP		

= 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 via an electronic form through the Jazz Safety tool of the Patient Support Program "ASOFARMA A TU LA" (Reference number: SV-ADIUM-SV-0014-20250718) on 18-Jul-2025 from a reporter (consumer or non-healthcare professional) regarding an elderly male patient who experienced serious (Death, life-threatening, disability/incapacity, hospitalization) 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 19-Jul-2025.

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

Concomitant medication was unknown.

On 27-Nov-2021, the patient began receiving Eligard 45 mg at an unknown frequency, via subcutaneous route, for prostate cancer (Lot numbers and Expiration dates were not provided).

On an unknown date in 2025, it was reported that the patient died 6 months ago. The cause of death was unknown. The patient was 74-year- 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, life-threatening, disability/incapacity, hospitalization).

The reporter assessed the causality of death in relationship to Eligard and Eligard Unspecified Device as related.

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

On 21-Jul-2025, the follow up was received from El Salvador by Adium via Patient Support Programme (Reference number: SV-ADIUM-SV-0014-20250718) and sent to Tolmar on 22-Jul-2025. New information included: dose frequency was added. All information was merged and processed together.

On 27-Nov-2021, the patient began receiving Eligard 45 mg every 6 months, via subcutaneous route, for prostate cancer (Lot numbers and Expiration dates were not provided).

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

## Listedness

Death >Eligard® >unlisted as per CCDS Eligard®> 7-Nov-2024

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

Death> Eligard®>unlisted as per USPI Eligard®>Feb-2025

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

## Company Remarks (Sender's Comments) :

Evaluator comment (Tolmar): This is regarding an elderly 74-year-old male patient who reported death (death) during Eligard (leuprolide acetate) 45 mg therapy for prostate cancer. Tolmar assessed death as serious due to fatal outcome and also life-threatening, disability/incapacity and hospitalization. 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

## Continuation Sheet for CIOMS report

Form of Admin : 1) Injection  
 Lot Number : 1) Unknown  
 Daily Dose : (45 milligram(s))  
 Route of Admin : 1) Subcutaneous  
 Indications : 1) prostate cancer [10060862 - Prostate cancer]  
 Therapy Dates : 1) From : 27/Nov/2021 To :Not applicable  
 Action(s) Taken With Drug : Not applicable

## Causality

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

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

## Labeling :

1) Death

CORE