													<u> </u>	<u> </u>		_	RIVI
SUSPECT ADVERSE REACTION REPORT																	
3031 E	JI ADVENSE I	KLACTION KLI O	IX I													_	
												Ш					
[				INFORM		1				_							
PATIENT INITIALS     (first, last)	1a. COUNTRY  COSTA RICA	2. DATE OF BIRTH  Day Month Year	2a. AGE 56	3. SEX	3a. WEIGHT Unk	Day		ACTION Month	_	ear	8-12	APF	ECK AL PROPR /ERSE	IATE		.1	
PRIVACY		PRIVACY	Years	Female				Unk				ADV	LKSL	NEA	CHO	V	
7 + 13 DESCRIBE REACTION(S) (including relevant tests/lab data) Event Verbatim [LOWER LEVEL TERM] (Related symptoms if any separated by commas)										PATIENT DIED							
Other Serious Criteria: Medically Significant									INVOLVED OR								
had developed a condition that left her with vision in only one eye [Blindness, one eye]											PROLONGED INPATIENT HOSPITALISATION						
Case Description: This is a spontaneous report received from a Consumer or other non HCP, Program ID:											_	LINIVA	OLVED	DEI	CICT	NIT	
164974.										INVOLVED PERSISTENT OR SIGNIFICANT DISABILITY OR							
A 56-year-old female patient received etanercept (ENBREL), (Batch/Lot number: unknown) at 50 mg weekly										y		INC	APACI	ΓΥ			
for arthritis. The patient's relevant medical history and concomitant medications were not reported.										_		_					
				(Contin	ued on Ad	dition	al In	formati	ion Pa	ige)	Ш	THE	REATEN	NING			
		II. SUSPEC	T DRUC	G(S) INF	ORMA	TIO	N										
II. SUSPECT DRUG(S) INFORMATION  14. SUSPECT DRUG(S) (include generic name)													ACTION AFTER		DDING	2	
#1 ) Enbrel (ETANERCEPT) Solution for injection in pre-filled syringe #2 ) Enbrel (ETANERCEPT (DEVICE CONSTITUENT)) Solution for injection in pre-filled syringe												RUG?	-11 ILIX	010	71 11 11 11	,	
15. DAILY DOSE(S)			16	. ROUTE(S) C	F ADMINIST		N				_	_					
#1 ) 50 mg, weekly #1											L	YES	3 <u> </u>	NO	Шм	IA	
#2 ) #2 ) Unknown 17. INDICATION(S) FOR USE												ACTION EAR AF					
#1 ) arthritis (Arthritis) #2 ) Unknown										R	EINTR	ODUC	TION	l?			
` ' '					THERAPY DURATION							٦٧,	. —		П.		
·					) Unknown :) Unknown						YES NO NA						
		III. CONCOMIT	ANTO	2110(0)	^ NID I I	IOT	<u> </u>	.,									
22. CONCOMITANT DRU	JG(S) AND DATES OF ADM	III. CONCOMIT		\ /	AND H	151	UK	ĭ									
23. OTHER RELEVANT I	HISTORY. (e.g. diagnostics	, allergies, pregnancy with last mo Type of History / Notes		etc.) Description													
Unknown		Type of History / Notes	'	Description													
		IV. MANUF	ACTUR	FR INF	ОВМАТ	ION	J										
24a. NAME AND ADDRE	26. REMA		101	1													
Pfizer S.A. Laura Arce Mora																	
Avenida Escazú, T San Jose, COST																	
	24b. MFR CC	ONTROL NO.		25b. NAM	E AND ADDR	RESS C	)F RF	PORTE	R								
		00098759		25b. NAME AND ADDRESS OF REPORTI NAME AND ADDRESS WITHH													
24c. DATE RECEIVED	24d, REPOR			$\dashv$													
BY MANUFACTURE	STUDY	LITERATURE															
13-AUG-2025	HEALTH		aneous	_													
DATE OF THIS REPORT 15-AUG-2025	25a. REPOR	T TYPE FOLLOWUP:															

## **ADDITIONAL INFORMATION**

## 7+13. DESCRIBE REACTION(S) continued

The following information was reported: BLINDNESS UNILATERAL (medically significant), outcome "unknown", described as "had developed a condition that left her with vision in only one eye". The action taken for etanercept was unknown. Clinical course: The patient not enrolled in the program indicated that she had been using the medication Enbrel for approximately nine years. She mentioned that, since the pandemic, she had developed a condition that left her with vision in only one eye. She was provided with the enrollment process so she could access the benefits. The patient stated that she suffers from arthritis and had been seen by several doctors. A meeting was held with multiple doctors because the one who had been treating them passed away.

The information on the batch/lot number for etanercept will be requested and submitted if and when received.