													CI	Oi	VIS	-01	RIVI
SUSPECT ADVERSE REACTION REPORT																	
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		I. F	REACTION	INFOR	MATION	I											
1. PATIENT INITIALS (first, last)	1a. COUNTRY	2. DATE OF BIRTI	H 2a. AGE	3. SEX	3a. WEIGHT	4-6	_	ACTION	_		8-12		ECK AL		: TO		
PRIVACY	PANAMA	PRIVACY	Year 40 Years	Female	Unk	Day		Month Unk		rear/			VERSE			1	
7 + 13 DESCRIBE REACTION(S) (including relevant tests/lab data) Event Verbatim [LOWER LEVEL TERM] (Related symptoms if any separated by commas) bruises were appearing on her skin, especially on her arms and in the belly area where ENBREL has been injected [Injection site bruising]											PATIENT DIED  INVOLVED OR PROLONGED INPATIENT HOSPITALISATION						
Case Description: This is a spontaneous report received from a Consumer or other non HCP, Program ID: 164974.  A 40-year-old female patient received etanercept (ENBREL), (Lot number: LL4200, Expiration Date: May2027)										027)	INVOLVED PERSISTENT OR SIGNIFICANT DISABILITY OR INCAPACITY						
at 50 mg weekly.  (Continued on Additional Information Page									age)	LIFE THREATENING							
II. SUSPECT DRUG(S) INFORMATION  14. SUSPECT DRUG(S) (include generic name) #1 ) Enbrel (ETANERCEPT) Solution for injection in pre-filled syringe {Lot # LL4200; Exp.Dt. MAY-2027} #2 ) Enbrel (ETANERCEPT (DEVICE CONSTITUENT)) Solution for injection in pre-filled syringe											20. DID REACTION ABATE AFTER STOPPING DRUG?						
15. DAILY DOSE(S) #1 ) 50 mg, weekly #2 )	#1 ) Unkno	ROUTE(S) OF ADMINISTRATION ) Unknown ) Unknown							YES NO NA								
17. INDICATION(S) FOR USE #1 ) Unknown #2 ) Unknown										21. DID REACTION REAPPEAR AFTER REINTRODUCTION?							
18. THERAPY DATES(from/to #1 ) Unknown #2 ) Unknown	#1 ) Unkno	THERAPY DURATION ) Unknown ) Unknown							YES NO NA								
#2 ) OHKHOWH				,		10.70	20				<u> </u>						
III. CONCOMITANT DRUG(S) AND HISTORY  22. CONCOMITANT DRUG(S) AND DATES OF ADMINISTRATION (exclude those used to treat reaction)																	
23. OTHER RELEVANT HIST From/To Dates Unknown	ORY. (e.g. diagnostics	allergies, pregnancy with Type of History / N		d, etc.) Description													
IV. MANUFACTURER INFORMATION  24a. NAME AND ADDRESS OF MANUFACTURER  26. REMARKS																	
Pfizer S.A. Laura Arce Mora Avenida Escazú, Torre San jose, COSTA R																	
	24b. MFR CC	ONTROL NO.		25b. NAME AND ADDRESS OF REPORTER NAME AND ADDRESS WITHHELD.													
24c. DATE RECEIVED BY MANUFACTURER	24d. REPOR		TIDE	$\neg$													
14-AUG-2025	STUDY HEALTH	LITERAT SSIONAL OTHER:	Spontaneous														
DATE OF THIS REPORT 15-AUG-2025	25a. REPOR	T TYPE	VUP:														

## **ADDITIONAL INFORMATION**

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

The patient's relevant medical history and concomitant medications were not reported.

The following information was reported: INJECTION SITE BRUISING (non-serious), outcome "unknown", described as "bruises were appearing on her skin, especially on her arms and in the belly area where ENBREL has been injected". The action taken for etanercept was unknown.

Additional information: The patient was feeling somewhat anxious and worried about the appearance of these bruises, and mentioned that she did not have an appointment with the rheumatologist until 30Sep2025.

Amendment: This follow-up report is being submitted to amend previous information: event data (removed: Bruise).