SUSPECT ADVERSE REACTION REPORT

CIOMS FORM

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SUSPECT AL	OVERSE REACTION	N REPC)RT									
I			I. REA	CTIC	ON INFOR	RMATIO	N					
1. PATIENT INITIALS	1a. COUNTRY	2. DATE OF BIRTH			2a. AGE	3. SEX	4-6 REACTION ON		ONSE	Γ	8-12 CHECK ALL	
(first, last)		Day	Month	Year	Years		Day	Month	n	Year		APPROPRIATE TO ADVERSE
7 + 13 DESCRIBE F	REACTION(S) (includi	ng relev	vant tests,	(Vab C	аата)							REACTION PATIENT DIED INVOLVED OR PROLONGED INPATIENT HOSPITALISATION INVOLVED PERSISTENT OR SIGNIFICANT DISABILITY OR INCAPACITY LIFE THREATENING
		II.	SUSPEC	T DF	RUG(S) IN	NFORM <i>A</i>	ATION					
ABA STC									ABAT STOP	ID REACTION E AFTER PPING DRUG? ES		
15. DAILY DOSE(S)					16. ROUTE(S) OF					21. D	ID REACTION	
					ADM	INISTRAT	ΓΙΟΝ			F	REAPI	PEAR
17. INDICATION(S) FOR USE							_	r reintro-duction? es $\ \square$ no $\ \square$ na				
18. THERAPY DATES (from/to) 19. THERAPY DURATION												
L		III. CC	DNCOMI	TAN	T DRUG(S) AND	HISTO	DRY				
22. CONCOMITANT DRUG(S) AND DATES OF ADMINISTRATION (exclude those used to treat reaction)												
23. OTHER RELEVANT HISTORY (e.g. diagnoses, allergies, pregnancy with last menstrual period, etc.)												

IV. MANUFACTURER INFORMATION

24a. NAME AND ADDRESS OF	MANUFACTURER	
	24b. MFR CONTROL NO.	
24c. DATE RECEIVED BY MANUFACTURER	24d. REPORT SOURCE STUDY LITERATURE HEALTH PROFESSIONAL	
DATE OF THIS REPORT	25a. REPORT TYPE INITIAL FOLLOW-UP	