CIOMS	FORM

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SUSPECT AD	VERSE REACT	ION REPORT																				
		I. REACTI	ON	INFOR	MATIC	 NC		1	1	ļ	.l									_		
1. PATIENT INITIALS	1a. COUNTRY	COUNTRY 2. DATE OF BIRTH				2a. AGE 3. SEX 4-6 REACTION ONSET											1 0 12 011EON ALL					
(first, last)		Day Month	Years Day Month Year										APPROPRIATE TO ADVERSE REACTION									
7 + 13 DESCRIBE	REACTION(S) (in	cluding relevant	tests	s/lab data	a)											ATIE			ED			
															PF IN	IVOI ROL: IPAT DSPI	ON:	GEE NT	)	l		
												i.			PE SIG DI:	IVOI RSIS GNIF SAB CAP	STEN FICA BILIT	NCE ANT Y C				
													□ LIFE THREATENING									
	II.	SUSPECT D	RUC	G(S) IN	FORM	IA٦	ΓΙΟΙ	N												_		
14. SUSPECT DRUG(	(S) (include gener	ic name)													AB ST	BATI OPP	E A	AFTE DR	ER UG?			
15. DAILY DOSE(S)				16. ROUTE(S) OF ADMINISTRATION							i	21. DID REACTION REAPPEAR										
17. INDICATION(S) F	OR USE												☐ AFTER REINTRO DUCTION? ☐ YES ☐ NO ☐ NA									
18. THERAPY DATES	(from/to)			19. TH	HERAPY	/ D	URA	TIO	N													
	III. Co	ONCOMITAN'	T D	RUG(S)	AND	Н	IST	OR	Υ													
22. CONCOMITANT	DRUG(S) AND DA	ATES OF ADMIN	IISTR	ATION (	exclude	e th	ose	use	ed 1	to t	tre	at r	ea	actic	on)							
23. OTHER RELEVAN	IT HISTORY (e.g.	diagnostics, alle	ergics	s, pregna	incy wit	ith	last	mo	nth	of	р	erio	d,	etc								
	IV	. MANUFAC	TUR	FR INF	ORMA	ΔΤ	ION															
24a. NAME AND ADI					2.1111/																	
	24b. M	FR CONTROL N	Ο.	-																		
24c. DATE RECEIVED BY MANUFACTU	JRER □ STU	PORT SOURCE  JDY																				
DATE OF THIS REPOF	i i	PORT TYPE	VUP																			