

SUSPECTED ADVERSE DRUG REACTION REPORTING FORM

For **VOLUNTARY** reporting of ADRs by Healthcare Professionals

INDIAN PHARMACOPOEIA COMMISSION (National Coordination Centre-Pharmacovigilance Programme of India)

Ministry of Health & Family Welfare, Government of India, Sector-23, Raj Nagar, Ghaziabad-201002

PvPI Helpline (Toll Free) :1800-180-3024 (9:00 AM to 5:30 PM, Monday-Friday)

Initial Case 🖬 Follow-up Case 🗖											FOR AMC / NCC USE ONLY									
A. PA	TIENT INFOR	MATI	ON *								Reg. No. / IPD No. / OPD No. / CR No. :									
1. Patient Initials: 2. Age or date of birth:											AMC Report No. :									
3. Gender: M 🗆 F 🗅 Other 🗅 4.Weight (in Kg.)										Worldwide Unique No. :										
											12. Relevant investigations with dates :									
B. S	USPECTED AD	VERS	E REACTIO	ON *	k															
5. Ev	vent / Reaction	start o	date (dd/m	m/y	ууу)															
	ent / Reaction		-																	
7. D	escribe Event/R	eactio	n managen	nent	with de	tails	, if ar	ıy												
			13. Relevant medical / medication history (e.g. allergies, pregnancy, addiction, hepatic, renal dysfunction etc.)																	
											pregnancy,	, ad	idiction, h	iepat	ic, ren	al dysf	unction etc	c.)		
											14. Seriousness of the reaction : No if Yes (<i>please tick anyone</i>)									
											Death (dd/mm/yyyy)									
			Life threa		-				sability											
			Hospitalization-Initial/Prolonged Other Medically important																	
			15. Outcome:																	
			Recovered Recovering Not Recovered Fatal Recovered with sequelae Unknown																	
C 6	ISPECTED ME	DICA	TON(S) *								□Fatal			Jvere	u with	seque		KIIU	VII	
S.	SUSPECTED MEDICATION(S) 8. Manufact			1 1			iry	Dose	F	Route	Frequency		Therap	ates	Inc	dication		Causality		
No.	Name		rer	1		Dat	te				,		Date	Date]		Assessment		
	(Brand/ Generic)	()	if known)	Lot No.		(ii knov						1	Started	Stoppe		*				
i																				
ii																				
iii																				
iv [#]												_								
9. Ad	tion taken afte	r reac	tion (<i>plea</i> s	e tic	:k)												after reintro (<i>please tic</i>		tion of	
S.	Drug		Dose		Dose		Dose not		Not		Unknown		Yes			0	Effect		Dose	
No.			ncreased		reduced		ange	d applic		able							unknov	vn	(if re-	
as per C																			introduced)	
i																				
ii																				
iii																				
iv																				
11.	Concomitant me	dical			-									de th	iose us	ed to				
Name Dose S. No. (Brand / Generic)					Rou	ite	Frequency (OD, BD, etc.)				Therapy	-	Dates				Indic	atio	n	
5. 10										Date Started			Date Stopped							
i																				
ii																				
iii [#]																				
Additional Information : D. R											PORTER DE	ETA	ILS *							
16. Na												ess	:							
											Email :									
											t No- :									
										-	ation :Signature :									
										17. D	ate of this	rep	ort (dd/	mm,	'yyyy)):				
	Signature and Name of Receiving Personnel : Confidentiality : The patient's identity is held in strict confidence a												6 . II							
Confi	dentiality : T	he pa	tient's id	enti	tv is he	ld in	stric	confid	ence	and pr	otected to t	the	fullest e	exter	it. Su	Dmiss	sion of a r	epo	rt does not	

constitute an admission that medical personnel or manufacturer or the product caused or contributed to the reaction. Submission of an ADR report does not have any legal implication on the reporter.

Use separate page for more information* Mandatory Fields for suspected ADR Reporting Form