

## SUSPECTED ADVERSE DRUG REACTION REPORTING FORM

For VOLUNTARY reporting of Adverse Drug Reactions by Healthcare Professionals

INDIAN PHARMACOPOEIA COMMISSION											FOR AMC/NCC USE ONLY										
(National Coordination Centre-Pharmacovigilance Programme of India) Ministry of Health & Family Welfare, Government of India Sector-23, Raj Nagar, Ghaziabad-201002											AMC Report No. :										
Report Type 🗆 Initial 🗆 Follow up											Worldwide Unique No. :										
A. PATIENT INFORMATION												12. Relevant tests/ laboratory data with dates									
1. Pa	. Patient Initials 2. Age at time Event or Date		Date o	of	3. N	И 🗆 F	🗆 Oth	er 🗆													
Birth																					
B. SUSPECTED ADVERSE REACTION												13. Relevant medical/ medication history (e.g. allergies, race,									
	5. Date of reaction started (dd/mm/yyyy)												pregnancy, smoking, alcohol use, hepatic/renal dysfunction etc.)								
	te of recov	-	(dd/m	ım/yy	уу)																
7. De	escribe reac	tion or																			
													14. Seriousness of the reaction: No $\Box$ if Yes $\Box$ (please tick anyone)								
											Death (dd/mm/yyyy)     Congenital-anomaly										
												□ Life threatening □ Required intervention to									
												Prevent permanent									
												Hospitalization/Prolonged impairment/damage									
						Disability Other (specify)															
												15. Outcomes         □ Recovered       □ Recovering       □ Not recovered									
													□ Fatal □ Recovered with sequelae □ Unknown								
C. SI	JSPECTED	MEDIO	CATION(S	)							Tu	-cui				in seque		CHRIGHT			
	C. SUSPECTED MEDICATION(S) 8. Name Manufacturer Batch No. Exp. Date Dose Route Freq											quency Therapy dates Caucality									
S.No	8. Name Manufacturer (Brand/Generic) (if known)			Batch No. Exp. Date Dose / Lot No. (if known) used				Route used	(OD, B	D				ed	Indication		Causality Assessment				
i										etc.)											
ii																					
iii																					
lv S No.	9 Action Ta	iken (nl	ease tick)							10 Rea	acti	on rea	nneare	d after rein	trodu	uction (nl	eace t	ick)			
as	9. Action Taken (please tick) Drug Drug Dose Not Unkn								0. Reaction reappeared after reintroduction (please tick)												
per C	withdrawn		ncreased	creased				nged applicable		Ye	es	s No		Effect unk		known Dose		(if reintroduced)			
i 									-												
ii iii																					
iv																					
11. C	Concomitan	t medic	al product	t inclu	iding sel	f-mec	dication	and her	bal rem	edies wit	h tł	nerapy	dates (	Exclude the	ose u	sed to tre	eat rea	action)			
S.No Name (Brand/Generic)					Dose u	ised		luency				py dates		Indication							
								(OD, BD,		etc.) Date:		tarted	Date stop	Date stopped							
i ii																					
iii																					
Additional Information: D. REF													REPORTER DETAILS								
16. N													Name and Professional Address:								
Pin:_													:E-mail								
Tel.												l. No. (with STD code)									
												cupation:Signature:									
														dd/mm/yy							
Confidentiality: The patient's identity is held in strict confidence and protected to the fullest extent. Programme staff is not expected to and will not disclose the reporter's identity in response to a request from the public. Submission of a report does not																					
	stitute an a																				

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Pharmacovigilance Programme of India for Assuring Drug Safety

### **ADVICE ABOUT REPORTING**

#### A. What to report

- > Report serious adverse drug reactions. A reaction is serious when the patient outcome is:
  - Death
  - Life-threatening
  - Hospitalization (initial or prolonged)
  - Disability (significant, persistent or permanent)
  - Congenital anomaly
  - Required intervention to prevent permanent impairment or damage
- Report non-serious, known or unknown, frequent or rare adverse drug reactions due to Medicines, Vaccines and Herbal products.

#### B. Who can report

> All healthcare professionals (Clinicians, Dentists, Pharmacists and Nurses) can report adverse drug reactions

#### C. Where to report

- Duly filled Suspected Adverse Drug Reaction Reporting Form can be send to the nearest Adverse Drug Reaction Monitoring Centre (AMC) or directly to the National Coordination Centre (NCC).
- > Call on Helpline (Toll Free) 1800 180 3024 to report ADRs.
- Or can directly mail this filled form to pvpi@ipcindia.net or pvpi.ipcindia@gmail.com
- > A list of nationwide AMCs is available at:

#### http://www.ipc.gov.in, http://www.ipc.gov.in/PvPI/pv\_home.html

#### D. What happens to the submitted information

- Information provided in this form is handled in strict confidence. The causality assessment is carried out at AMCs by using WHO-UMC scale. The analyzed forms are forwarded to the NCC through ADR database. Finally the data is analyzed and forwarded to the Global Pharmacovigilance Database managed by WHO Uppsala Monitoring Centre in Sweden.
- The reports are periodically reviewed by the NCC-PvPI. The information generated on the basis of these reports helps in continuous assessment of the benefit-risk ratio of medicines.
- The information is submitted to the Steering committee of PvPI constituted by the Ministry of Health & Family Welfare. The Committee is entrusted with the responsibility to review the data and suggest any interventions that may be required.

#### E. Mandatory field for suspected ADR reporting form

Patient initials, age at onset of reaction, reaction term(s), date of onset of reaction, suspected medication(s) and reporter information.

For ADRs Reporting Call on PvPI Helpline (Toll Free)

# 1800 180 3024

(9:00 AM to 5:30 PM, Working Days)