SUSPECTED ADVERSE DRUG REACTION REPORTING FORM

For VOLUNTARY reporting of Adverse Drug Reactions by Healthcare Professionals

NEON LABORATORIES LIMITED 140 Damji Shamji Ind.Complex,M.Caves Road.,											ica		ricaltine	larc	110103310114	NEON		
Andheri (E),Mumbai-400093. Ph:+918767559938											Case Id :							
Email: drugsafety@neongroup.com											(To be filled by Neon laboratories Ltd)							
Report Type Initial Follow up																		
A. PATIENT INFORMATION												12. Relevant tests/ laboratory data with dates						
1. P	atient Initial		2. Age a			3. M \Box F \Box Other \Box												
			Event or Date Birth		-	4. Weight Kgs												
B. SUSPECTED ADVERSE REACTION								I	ngs	13	13. Relevant medical/ medication history (e.g. allergies, race,							
							pregnancy, smoking, alcohol use, hepatic/renal dysfunction etc.)											
	ate of reacti																	
	ate of recov escribe reac	(dd/m	m/yyy	yy)														
/. D	escribe reac		JIODIEIII							14								
												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											
						15.	15. Outcomes											
							□ Recovered □ Recovering □ Not recovered											
								\Box Fatal \Box Recovered with sequelae \Box Unknown										
C. 5	SUSPECTE	ED ME	DICATI	ON(S	5)													
Sr.	8. Name (Brand/Generic)		Manufacturer (if known)		Batch N / Lot N			te Dose used	Route used	Frequence (OD, BE		The Date	Therapy dates		Indicatio	n Causality Assessment		
No					`		own)	useu		etc.)		started						
i																		
ii iii																		
Iv																		
Sr.	9. Action T	aken (pl	ease tick))				- I		10. Read	ctio	on reappea	red afte	r re	introduction	(please tick)		
No as	Drug withdrawn Dose in		neresced		Dose duced	Dose not changed			Unkn own	Ye	es		No			Dose (if reintroduced)		
per C							-	<u> </u>							unknown			
i ii														-				
iii																		
iv																		
	-		-	t inclu							the	* *				treat reaction)		
Sr. No					Dose used Route used					uency BD,	The Date starte		erapy dates		ite stopped	Indication		
INO									etc.)	DD,		Jale started	а D		ite stopped			
i 													-					
ii iii																		
Additional Information:											D. REPORTER DETAILS							
						-	Name and Professional Address:											
											in:E-mail el. No. (with STD code)							
											Decupation:Signature:							
										17. Date	7. Date of this report (dd/mm/yyyy):							

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 Neon laboratories Ltd.
- > Call on Helpline +918767559938 to report ADRs.
- > Or can directly mail this filled form to drugsafety@neongroup.com
- A list of nationwide AMCs is available at: http://www.ipc.gov.in, http://www.ipc.gov.in/PvPI/pv_home.html

D.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 Helpline number

