


SUSPECTED ADVERSE DRUG REACTION REPORTING FORM

For VOLUNTARY reporting of Adverse Drug Reactions by Healthcare Professionals

Pharmacovigilance Operations Scientific Department Mapra Laboratories Private Limited										FOR MAPRA USE ONLY			
 <p style="font-size: small;">Group Company of ARISTO Pharmaceuticals Private Limited 201, Adhyaru Indl. Estate, Sun Mill Compound, Lower Parel, Mumbai - 400 013, Maharashtra, India. E-Mail: aepvc.scientific@mapra.com Toll free No. 1800225960 (Monday to Friday between 9.30 am to 5.30 pm, except on public holidays), WhatsApp: +918879607724.</p>										Report No.:			
A. PATIENT INFORMATION										Report Type: <input type="checkbox"/> Initial <input type="checkbox"/> Follow up			
1. Patient Initials _____		2. Age at time of Event or Date of Birth _____		3. M <input type="checkbox"/> F <input type="checkbox"/> Other <input type="checkbox"/>		Worldwide Unique No.:							
		4. Weight _____ Kgs		12. Relevant tests/laboratory data with dates									
B. SUSPECTED ADVERSE REACTION													
5. Event/Reaction start date (dd/mm/yyyy)					13. Relevant medical/medication history (e.g. allergies, race, pregnancy, smoking, alcohol use, hepatic/renal dysfunction, past surgery etc.)								
6. Event/Reaction stop date (dd/mm/yyyy)													
6 (A). Onset Lag Time													
7. Describe Event/Reaction with treatment details, if any													
					14. Seriousness of the reaction: No <input type="checkbox"/> if Yes <input type="checkbox"/> (please ck anyone)								
					<input type="checkbox"/> Death (dd/mm/yyyy)		<input type="checkbox"/> Congenital anomaly						
<input type="checkbox"/> Life threatening		<input type="checkbox"/> Disability											
<input type="checkbox"/> Hospitalization / Prolonged		<input type="checkbox"/> Other Medically important											
					15. Outcomes								
					<input type="checkbox"/> Recovered		<input type="checkbox"/> Recovering		<input type="checkbox"/> Not recovered				
<input type="checkbox"/> Fatal		<input type="checkbox"/> Recovered with sequelae		<input type="checkbox"/> Unknown									
C. SUSPECTED MEDICATION(S)													
S.No	8. Name (Brand/Generic)	Manufacturer (if known)	Batch No. / Lot No.	Exp. Date (if known)	Dose used	Route used	Frequency (OD, BD etc.)	Therapy dates		Indication	Causality Assessment		
								Date started	Date stopped				
i													
ii													
iii													
iv*													
S.No as per C	9. Action Taken (please tick)						10. Reaction reappeared after reintroduction (please tick)						
	Drug withdrawn	Dose increased	Dose reduced	Dose not changed	Not applicable	Unkn own	Yes	No	Effect unknown	Dose (if reintroduced)			
i													
ii													
iii													
iv													
11. Concomitant medical product including self-medication and herbal remedies with therapy dates (Exclude those used to treat reaction)													
S.No	Name (Brand/Generic)	Dose used	Route used	Frequency (OD, BD, etc.)	Therapy dates		Indication						
					Date started	Date stopped							
i													
ii													
iii*													
Additional Information:						D. REPORTER DETAILS							
						16. Name and Professional Address: _____							
						Pin: _____ E-mail _____							
						Tel. No. (with STD code) _____							
						Occupation: _____ Signature: _____							
						17. Date of this report (dd/mm/yyyy): _____							
Sign. and Name of Receiver-													
Confidentiality: The patient's identity is held in strict confidence and protected to the fullest extent. Submission of a report 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													