🙏 Paviour	PAVIOUR PHARMACEUTICALS PVT. LTD.							
Obsessed with Quality	PHARMACOVIGILANCE DEPARTMENT							
ADVERSE EVENT (AE) REPORT FORM								
Report Type:	J Initial	Follow-up		Follow-up No	D:			
Date of AE Report:								
Please fill and return this form to Paviour Pharmaceuticals Pvt. Ltd. 311-312, Suneja Tower-1,								
District Centre, Janak Puri, New Delhi - 110058 within 24 hours of knowledge of adverse event E-mail ID: drugsafety@paviour.org								
1. Patient Information		<u>norg</u>						
Initials/identifier:		Date of Birth (e.g. 01 Jan	Eth	Ethnic Origin:				
		1940)		□ White □ Asian □ Black/African				
			American 🗖 Other, Please Specify					
Sex: Male Female		Height (cm):	Weight (kg):					
Pregnant: [] Yes [] No		Country of occurrence:	Tel. No:					
2. Adverse Event In	formatio	n						
AE term(s):								
Course of event:								
Onset of AE or date	and time	when AE occurred:		Date:	Time:			
Onset of AE or date applicable:	and time	when event became serious, if		Date:	Time:			
Present Status:								
\Box Ongoing \rightarrow AE currently treated \Box Yes \Box No								
Resolved Please Specify Date:Time:								
Case description: Detailed description of the event (Include related signs/symptoms, course, outcome)								
Reason for seriousness:								
□ resulted in death □ life-threatening □ required inpatient hospitalization or prolongation of existing hospitalization □ resulted in persistent or significant disability/ incapability (as per reporter's								
opinion)/ congenital anomaly/ birth defect other medically important event (reporter's discretion)								
Intensity: Mild Moderate Severe								
Reporter's Causality: [] certainly [] probably [] possibly [] unlikely [] conditional [] unassessable [] not related								
Outcome of AE:								
Completely recovered/resolved Ongoing Fatal Lost to follow-up								
\Box Unknown \Box Recovered with sequelae \rightarrow specify:								

If outcome is f											
Cause of death											
Report of Autopsy available? No Yes (Please attach copy to this report)											
Further information:											
Lab test Details (with dates, results and normal range):											
3. Drug Details											
Name of the drug: Strength: Indication:											
Route of Adm	of Admin:Dosage form:Dose:										
Frequency: Expiry date:											
Start date:			Stop da	te:		Ongoing:					
Action taken w	vith sus	pect drug	g:								
 Dosage changed temporarily: Date:O Dosage reduced ODosage increased Drug stop temporarily: Date: 											
O Drug restarte	•	-									
Drug withdra	awn pern	nanently (Dosage not o	changed 🗖	Unknown	Not appli	cable				
Additional sus	spect dr	ug (if any	/) details as at	oove:							
	Event abated after drug Event reappeared after If yes, did reaction recur? stopped or dose reduced: reintroduction of suspect drug:						eaction recur?				
						🗆 Yes 🗖 No					
□ Not applicable		□ Not applicable			Not applicable						
4. Patient's Relevant Medical History (Supplement attached Yes/No)											
(E.g. concomitant diseases, previous history of present condition, allergy, drug or alcohol abuse)											
5. Concomit Drug Name	Dose	gs Route	Frequency	Start	Stop	Ongoing	Causal				
(generic)	/ Unit	Noute	Trequency	date	date	Ongoing	relationship to event				
							NonePossible				
Indication:											
							NonePossible				
Indication:											
							□ None □ Possible				

6. Reporter Details				
Name: Address: Country: Tel. No: Email:	Occupation: [] Physician [] Pharmacist [] Nurse [] Consumer [] Other, specify: Also reported to: [] Regulatory Authority [] Distributor [] None Date : , Signature:			
7. Send this report to:	8. To be filled by Manufacturer:			
Paviour Pharmaceuticals Pvt. Ltd. 311-312, Suneja Tower-1, District Centre, Janak Puri, New Delhi – 110058 Tel No. +91-11-25542181 E-mail: <u>drugsafety@paviour.org</u>	Date received by receiver: Name and sign of receiver: Safety Report ID:			