Format No.: MUM/CPC/002/2011/F-01-R00

	Ipca Laboratories Limited, Mumbai Corporate Pharmacovigilance Cell ADVERSE EVENT REPORTING FORM (In Confidence)	S kipca
PATIENT INFORMATION 1. Patient Initials : 2. Country :	3. Sex : M F 4. Age at time of event : years OR 5. Date of Birth: (dd/mm/yyyy)	6. Weight : Kg
ADVERSE EVENT 1. Do you consider the adverse even 2. If yes, please indicate why the adv Death	verse event is considered to be serious : (Check all that apply) Disability or Permanent damage Life-threatening Olonged Other important medical events	Congenital anomaly / birth defect
 If the adverse event is not serious Date of onset of event:	d hospital discharge summaries as required) , indicate intensity of the adverse reaction : Mild Mild (dd/mm/yyyy) 5. If event stopped, date: Mild (hh/mm) Time (if available) : of reaction(s), including body site and severity as well as description pecific diagnosis for the reaction)	Moderate Severe (dd/mm/yyyy) (hh/mm) on of signs and symptoms.
7. Information on recovery and ar Other:	Not recovered Fatal	Recovered with sequelae
 Setting where event occurred: Hospital Out-Patient Relevant tests / laboratory data, ir 		
10. Other relevant history, including pr hepatic / renal dysfunction, etc.	reexisting medical conditions: (e.g. allergies, race, pregnancy, smoking a	ind alcohol use,
11. Treatment of adverse event :		

#1	. Generic Name 2. Brand Name			3. Dosage form & labeled strength		4. Lot No./Batch No.		5. Expiry Date	6. Manufacturer
π I									
# 2									
# 3									
# 4	<u> </u>								
7. Daily Dose (Specify units - mg, ml, mg/kg) 8. Frequency			9. Route of administration		10. Indication for use of suspected drug		11. Therapy dates (if unknown, give duration Start End		
# 1							5	(dd/mm/yyyy)	dd/mm/yyyy)
# 2								Therapy duration:	: (days)
								Therapy duration:	
# 3								(dd/mm/yyyy) Therapy duration:	(dd/mm/yyyy) (days)
# 4								(dd/mm/yyyy) Therapy duration:	(dd/mm/yyyy) (days)
12. Event abated	after use s	stopped or	dose reduced	:	13 Fv	ent re	appeared	after reintroduction :	
#1 Yes				-	Yes		No	Not applicab	
#2 Yes	No		pplicable		Yes		No	Not applicab	_ <u></u>
Generic name # 1		form and strength	(Specify unit: e.g. mg, ml, mg	s- _{//kg)} adminis		use c	of the drug	Starting date	Stopping date
# 2								(dd/mm/yyyy)	(dd/mm/yyyy)
# 3								(dd/mm/yyyy)	dd/mm/yyyy)
	1								1
# 4								(dd/mm/yyyy)	(dd/mm/yyyy)
REPORTER	ess							(dd/mm/yyyy)	(dd/mm/yyyy)
REPORTER . Name and addre								(dd/mm/yyyy)	(dd/mm/yyyy)
REPORTER	Code)		3. +	lealth profess					
# 4 REPORTER 1. Name and addre Fel. No. (With STD 0 2. Date of this Repo 5. Also rported to :	Code) ort	dd/mm/yyy	y)	lealth profess		Yes	No		
REPORTER 1. Name and addre Tel. No. (With STD (2. Date of this Repo	Code) ort	dd/mm/yyy ory author	y) ity Distr	lealth profess	ional?	Yes	No] 4. Occupation _	
REPORTER 1. Name and addre Tel. No. (With STD 0 2. Date of this Report 5. Also rported to : Signature	Code) ort(dd/mm/yyy ory author	y) ity Distr	lealth profess	ional?	Yes	No] 4. Occupation _	
REPORTER 1. Name and addre Fel. No. (With STD 0 2. Date of this Repo 5. Also rported to :	Code) ort (r Regulato transform to: covigilance Ltd. dustrial Es umbai 400 0	e Cell tate 067, India	3. F	lealth profess	ional?	Yes	No] 4. Occupation _	