

Adverse Event Form

Patient Initials 2. Age at time of event 3. Gender or M F Date of Birth . Weight: kg. dd / mm / yyyy S. Adverse Event	 14. Other relevant history, including pre-existing medical conditions (e.g., allergies, race, pregnancy, smoking alcohol use, hepatic/renal dysfunction, etc.)
Date of Birth	race, pregnancy, smoking alcohol use, hepatic/renal
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. Weight:// 5. Country:kg. dd / mm / yyyy	race, pregnancy, smoking alcohol use, hepatic/renal
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. Adverse Event	
Adverse Event	
. Date when event started (dd/mm/yyyy)://	15. Seriousness of the event: Is the event Serious or Non-seriou
	If Serious, then Choose the criteria
. Date of recovery (dd/mm/yyyy):/	Death (dd/mm/yyyy)/_/ Results in persistent or significant disability/incapacity
. Describe event:	Life threatening
	Other medically important condition
	16. Outcomes
. Relationship of event to the suspected medication	Fatal Continuing Recovering Recovered
Related Not related	Other (specify)
C. Suspected medication(s) Name (brand and / or generic Manufacturer Sr.No. name) (if known) Batch No. / Lot No. (if known)	Exp. Date Dose Route Date Date use or pres- cribed for (if known) used Frequency started stopped cribed for
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ii iii	
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11. Action taken with respect to Suspect Drug	
None Dose reduction	Date of dose reduction :/ (dd/mm/yyyy)
Drug temporarily discontinued Date stopped :	_// (dd/mm/yyyy) Date re-started :/ (dd/mm/yyyy)
Drug permanently discontinued Date stopped :	_// (dd/mm/yyyy)
Dose increased Date of dose increased	sed :/ (dd/mm/yyyy)
12. Concomitant medical products and therapy dates including self-medication and herbal	Reporter Healthcare Professional Consumer Othe
	Name and Address:

Pin code:	E-mail:
Cell No./Tel. No. with STD Code:	
Specialty:	Signature:
18. Occupation:	19. Date of this report (dd/mm/yyyy)
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E. Consent	If Report is from a Consumer, Permission Given to Follow Up with HCP? Yes No HCP Contact Details
Tel. no.: 18002677779	email ID.: drugsafety@cipla.com