## A. PATIENT INFORMATION

1. Patient Initials: $\square$
2. Country: $\qquad$
3. Sex:

4. Age at time of event :
 year OR
5. Date of Birth
 (dd/mm/yyyy)
6. Weight :
 Kg

## B. ADVERSE EVENT

1. Do you consider the adverse event to be serious? $\square$ Yes
 No
2. If yes, please indicate why the adverse event is considered to be serious : (Check all that apply)

$\square$ Hospitalization - initial or prolonged $\square$ Other important medical events

If patient died, cause of death and post mortem findings: $\qquad$
$\qquad$
(Please attach autopsy findings and hospital discharge summaries as required)
3. If the adverse event is not serious, indicate intensity of the adverse reaction :
 Mild
 Moderate
 Severe
4. Date of onset of event:
 (dd/mm/yyyy)
5. If event stopped, date:
 (dd/mm/yyyy) Time (if available)
 (hh/mm) Time (if available) $\square$ : $\square$ (hh/mm)
6. Describe event: (Full description of reaction(s), including body site and severity as well as description of signs and symptoms. Whenever possible, describe a specific diagnosis for the reaction)
$\qquad$
$\qquad$
. Information on recovery and any sequelae:


Recovered with sequelae
Unknown $\quad \square$
Other: (

$\qquad$
$\qquad$
$\qquad$
8. Setting where event occurred:

Hospital $\square$ Out-Patient $\square$ Home $\square$ Nursing Home $\square$
9. Relevant tests / laboratory data, including dates:
$\qquad$
$\qquad$
$\qquad$
10. Other relevant history, including preexisting medical conditions: (e.g. allergies, race, pregnancy, smoking and alcohol use, hepatic / renal dysfunction, etc.
11. Treatment of adverse event:

## C. SUSPECT MEDICATION(S)


12. Event abated after use stopped or dose reduced :

13. Event reappeared after reintroduction:
Related $\square$
14. Relationship of the adverse event with the drug: (Please tick in appropriate box)
15. Concomitant medical products and therapy dates including self medication and herbal remedies: (exclude those used to treat the event)

| Generic name | Dosage form and Labeled strength | Daily dose (Specify unitse.g. $\mathrm{mg}, \mathrm{ml}, \mathrm{mg} / \mathrm{kg}$ ) | Route of administration | Indication for use of the drug | Starting date | Stopping date |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| \# 1 |  |  |  |  |  | $\square$  <br> (dd/mm/yyyy)  |
| \# 2 |  |  |  |  |  | $\square$  <br> (dd/mm/yyyy)  |
| \# 3 |  |  |  |  |  |  |
| \# 4 |  |  |  |  |  | $\square$  <br> (dd/mm/yyyy)  |

## D. REPORTER

1. Name and address $\qquad$
$\qquad$

Tel. No. (With STD Code)
2. Date of this Report $\square$ 3. Health professional? Yes
 No $\square$ 4. Occupation $\qquad$ (dd/mm/yyyy)
5. Also rported to: Regulatory authority $\qquad$ Distributor $\square$ No one else $\square$
6. Report Type: Initilal

Follow-up

## Signature

## Please send this form to:

Corporate Pharmacovigilance Cell

## Ipca Laboratories Ltd.

142-AB, Kandivli Industrial Estate
Kandivli (West), Mumbai 400 067, India
T: +91 2266474105
E: pharmacovigilance.mumbai@ipca.com

## If any additional data, then attach with this form

