



**7. List of other drugs used by the patient during the period of one month:**

Name of the drug	Manufacturer / Batch no.	Dose	Form / Route of administration	Date of		Reason for use	Any unwanted occurrences
				Starting	Stopped / Continued		

**8. Details of the drug suspected to cause ADR:**

- Name of the drug:
- Manufacturing date and Expiry date (if available):
- Remaining pack / label (if available):
- Consumed orally along with (water / milk / honey / or any other)
- Whether any dietary precautions have been prescribed?  
If yes, please specify:
- Whether the drug is consumed under medical supervision or used as self medication.
- Any other relevant information associated with drug use:

**9. Management provided / taken for suspected adverse reaction**

**10. Please indicate outcome of the suspected adverse reaction (tick appropriate)**

Recovered:	Not recovered:	Unknown:	Fatal:	If Fatal Date of death:
Severe: Yes / No.	Reaction abated after drug stopped or dose reduced:			
	Reaction reappeared after re administration of drug:			
Was the patient admitted to hospital? If yes, give name and address of hospital				

**11. Any abnormal findings of relevant laboratory investigations related to the episode done pre and post episode of ADR:**

## 12. Particulars of ADR Reporter:

<b>Please tick:</b> Patient / Attendant / Nurse / Doctor / Pharmacist / Health worker / Drug Manufacturer / Any others (please specify)
<b>Name:</b>
<b>Address:</b>
<b>Telephone / E - mail:</b>

Signature of the reporter:

Date:

**Please send the completed form to: The centre from where the form is received or to**

The Coordinator, National Pharmacovigilance Coordination Centre (NPvCC)

All India Institute of Ayurveda (AIIA), Mathura Road, Gautam Puri,

Sarita Vihar, New Delhi - 110 076

E-mail: [pharmacovigilanceayush@gmail.com](mailto:pharmacovigilanceayush@gmail.com), [ayush-pharmavig@aiia.gov.in](mailto:ayush-pharmavig@aiia.gov.in)

### The ADR Probability Scale

(Program Coordinator has to fill this scale)

	Questions	Yes	No	Don't Know
1	Are there previous conclusive reports on the reactions?	+1	0	0
2	Did the ADR appear after the suspected drug was administered?	+2	-1	0
3	Did the ADR improve when the drug was discontinued a specific antagonist was administered?	+1	0	0
4	Did the adverse reaction reappear when the drug was re-administered?	+2	-1	0
5	Are there alternatives causes that could solely have caused the ADR?	-1	+2	0
6	Was the drug detected in the blood (or other fluids) in a concentration known to be toxic?	+1	0	0
7	Was the reaction more severe when the dose was increased, or less severe when the dose was decreased?	+1	0	0
8	Did the patient have a similar reaction to the same or similar drugs in any previous exposure?	+1	0	0
9	Was the adverse event confirmed by objective evidence?	+1	0	0
	<b>Total Score</b>			

**Score: > 9 = Certain;**

**5-8 = Probable;**

**1-4 = Possible;**

**0 = Unlikely**

The Suspected Adverse Event	Grade - 1 (Mild)	
	Grade - 2 (Moderate)	
	Grade - 3 (Severe)	
	Grade - 4 (Threatening)	
The Suspected Adverse Event	Serious	
	Non-Serious	
The Suspected Adverse Event is due to	Physician	
	Patient	
	Drug	
	Other factors*	

Signature  
**Program Coordinator**