Reporting form for Suspected Adverse Reactions NATIONAL PHARMACOVIGILANCE PROGRAM AYURVEDA, SIDDHA, UNANI and HOMOEOPATHY (ASU & H) DRUGS

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1.4			

Personal information of the consumers / patients / ADR reporter's will be kept confidential.

AYURVEDA

vC ITRA, JAMNAGAR

Code [000]

Ay-ITRA/000/ /2023

All suspected reactions are to be reported with relevant details.

1.	Patient /	consumer identification	(please com	plete or tick	boxes below	as appropriate)

Name	Patient Record Number (PRN)			
Place of Birth	IPD / OPD			
Address:		Age:		
Village / Town		Sex: Male / Female		
Post / Via				
District / State				
Diagnosis:	Constitution and Temperar	Constitution and Temperament:		
	-			

2. Description of the suspected Adverse Reactions

	The transfer of the transfer o				
Date and time of initial observation					
Description of reaction					

3. Whether the patient is suffering with any chronic disorders?

Hepatic Renal Cardiac Diabetes Any Others (Specify, if others)

- **4. Addictions, if any?** If yes, please specify:
- **5.** H/O previous allergies / Drug reactions, if any: If yes, please specify:

6. List of all ASU & H drugs used by the patient during the period of one month:

	Manufacturer / Batch no. Dose	Manufacturar	For	Form / Route of	Dat	te of	Reason	Any unwanted
Name of the drug		LINCA	administration	Starting	Stopped / Continued	for use	occurrences	

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

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

8. Details of the drug suspected to cause ADR:

- a. Name of the drug:
- b. Manufacturing date and Expiry date (if available):
- c. Remaining pack / label (if available):
- d. Consumed orally along with (water / milk / honey / or any other):
- e. Whether any dietary precautions have been prescribed? If yes, please specify:
- f. Whether the drug is consumed under medical supervision or used as self medication:
- g. 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 a	Reaction reappeared after re administration of drug:			
Was the patient admitted to hospital? If					
ves, 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:

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

Signature of the reporter:

Date:

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

To The Coordinator,

Intermediary Pharmacovigilance Centre for Ayurveda

Institute of Teaching and Research in Ayurveda, Jamnagar, Gujarat – 361008, India Tele Fax: 0288 2676856 / 0288 2553936

 $Website: \underline{https://itra.ac.in/}(\ hospital,pharmacovigilance)\ , Email: \underline{ipvcjamnagar@gmail.com}$

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
	Total Score			

Score: > 9 = Certain; 5-8 = Probable; 1-4 = Possible; 0 = Unlikely

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

If possible, explain the other factors: