

# REPORT ON SUSPECTED ADVERSE DRUG REACTIONS

#### NATIONAL CENTRE FOR ADVERSE DRUG REACTIONS MONITORING

Email: fv@npra.gov.my Website: www.npra.gov.my (Please report all suspected adverse drug reactions including those for vaccines, health supplements and traditional products. Do not hesitate to report if some details are not known. Mandatory fields are marked with \*, but please give as much other information as you can. Identities of Reporter, Patient and Institution will remain Confidential.)

	REPORT No. (for of	ticiai use oniy)	:			
PATIENT INFORMATION						
I.C. No. / R/N / Initials	*Age *Ge Male	ender <i>(please tid</i> Female		Wt (kg)	*Ethnic G	roup    Please tick (if applicable):   Initial Report   Follow-up Report
*ADVERSE REACTION RESCRIPT	FION /inc. company	of advance ov	outo dotailo d	of we also also as	intonation	
*ADVERSE REACTION DESCRIPT	IION (Inc. sequence	or adverse ev	ents, details d	or rechallenge	e, interaction	S)
						D
Time to onset of reaction :	,	Date start of reaction : Date end of reaction :				
	(please circle)					
Reaction subsided after stopping drug / reducing dose : Yes No Unknown *N/A (drug continued)						
Reaction reappeared after reintroducing drug : Yes No Unknown M/A (not reintroduced)						
Extent of reaction : Mild Moderate Severe						
Seriousness Life	Caused or prolon	-	Caused disab	oility	Caused bir	th
of reaction : threatening	hospitalisation		or incapacity		defect	(not serious)
Treatment of adverse reaction & action taken :						
Outcome : Recovered Recovering Not Victoria Vict						
Drug-reaction relationship : Certain Probable Possible Unlikely Unclassifiable						
*Suspected Drug(s):						
	Dose &		Batch / Lot	Therap	v Dates	
Product / Generic Name	Frequency Given	MAL No.	No.	Start	Stop	Indication
	Given					
	(.1	. / Ond / Ord / I		<u> </u>	D:1	1. D. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1.
For Vaccines Only: Vaccine dose (please circle): 1st / 2nd / 3rd / booster/ others: Diluent Batch / Lot No.:						
Concomitant Drug(s) / Other Vaco	cine(s) given just p	rior to AEFI [a	dverse events f	following immu	nisation] <i>(ple</i>	ase state 'NIL' if none):
Draduct / Canadia Nama	Dose &		Batch / Lot	Therap	y Dates	lu dia atia u
Product / Generic Name	Frequency Given	MAL No.	No.	Start	Stop	Indication
(Please attach additional sheets if r	necessary)					
(Frease allacif additional sheets if F	iecessary)					
Relevant Investigations	s / Laboratory Data		, ,		vant Medic	
· ·	•		(e.g.: nepa	tic / renal dys	stunction, alle	ergies, pregnancy status, etc)
Reporter Details						
*Name :	*1504:4	tion Nama				
INAITIC .	ame : *Institution Name					
Designation :	*Tel No	n ·				
*Email Address :	Date of Report :		Signature :			revision-01

## **ADR Reporting Guide**

Before submitting your ADR report, do check if you have inserted the following information.

\*Please try to fill every section in the ADR form overleaf, stating 'none / nil' if applicable. A complete report is a useful report.

#### NO. IMPORTANT POINTS TO NOTE

- 1 Definitions:
  - (i) Time to onset of reaction: time interval between first dose (initiation) of the drug until first sign of the ADR.
  - (ii) Initial report: First submission of report to NPRA of a particular patient involving a particular ADR.
  - (iii) Follow-up report: Submission of further reports related to the same case to inform of additional information not mentioned previously or which occurred after the initial report. Please mention the date of initial report for reference.
- 2 Please specify any previous history of allergy (including drugs, food, etc.).
- 3 Include information on any concomitant medications or underlying illnesses? (Please state 'nil' if none)
  - Date started and stopped for each medication
  - · Please state 'cont' for any medication still continued after the ADR
- 4 Please state the specific **indication** of the suspected drug (e.g.: 'pneumonia due to S. Pneumoniae' not 'infection' or 'antibiotic').
- If the ADR reappeared after reintroducing drug (rechallenge), please describe the rechallenge fully (dose given, timing, brand used, etc.) under section 'Adverse Reaction Description'.
- 6 Please specify if any treatment was given for the ADR, or if the suspected drug was stopped, what alternative drug was started and how the patient responded.
- 7 Please include the latest / current **outcome** of the patient (e.g. recovered fully, not recovered).
  - If possible, follow-up the patient periodically until the final outcome is known.
  - A follow-up report may be sent in to update on the final outcome of the patient.
- 8 **Skin reactions**: Please describe the specific type and location of the skin reaction. (Use the Cutaneous ADR form and guide available on www.npra.gov.my)
- 9 Do keep your own record of details enabling you to **contact** the patient or trace the case notes later on if necessary (e.g. IC number, patient name and phone number).

Please refer to our website for additional guidance on ADR Reporting, or contact us at fv@npra.gov.my if you have any queries.

### Laporan Kesan Advers Ubat

Bahagian Regulatori Farmasi Negara (NPRA) Kementerian Kesihatan Malaysia

PUSAT PEMONITORAN KESAN ADVERS UBAT KEBANGSAAN
BAHAGIAN REGULATORI FARMASI NEGARA
LOT 36, JALAN UNIVERSITI
46200 PETALING JAYA
SELANGOR