

For office use only:				
BRN				
Initial report				
Follow-up report				

National Adverse Drug Reaction Monitoring Centre Ministry of Health, Brunei Darussalam

SUSPECTED ADVERSE DRUG REACTION REPORT FORM

Please report **all** suspected adverse drug reactions including those for self-medication, traditional medicines and health supplements. Do not hesitate to report if some details are not known. **MANDATORY FIELDS** are marked with *. **Identities of reporter and patient** will be kept confidential.

(1) PATIENT DETAILS *

Patient name:			I	Date of birth:	Weight,	if known (kg):
Gender: 🗌 Male 🗍 Female Medical record no. / BruHims no. / Identity card no. :						
Nationality: Ethnic group: 🗖 Malay 🗍 Chinese 🗖 Other (please specify):						
(2) ADVERSE DRUG REACTION (ADR) DETAILS *						
Description of ADR(s):						
Time to onset of ADR(s):mins/ hours/ days/ months/ years Date ADR(s) started:Date ADR (s) stopped: (please circle)						
Do you consider the ADR(s) to be seriou	ıs? 🗆 Yes 🛛 No				
If yes , please indicate	e why the AI	OR is considered to be s	erious (please tick all the	at apply):		
 Patient died due to reaction Life threatening Involved or prolonged in-patient hospitalisation Congenital abnormality Involved persistent or significant disability or incapacity Medically significant; please give details:						
Description of treatment						
						s or injuries)
Outcome of reaction: Recovered Recovering Recovered with sequelae (any permanent complications or injuries) Not recovered Fatal (Date of Death): Unknown						
(3) SUSPECTED DRUG(S) * (Additional pages may be attached)						
Product name, active	Dosage	Dosage regimen	Product details:	Date	Date	Prescribed for
ingredient & strength	form	(dosage, frequency & route)	Batch No. (if known)	started	stopped	1105011504101
1.						
2.						
3.						
(4) OTHER DRUG(S) (INCLUDING SELF-MEDICATION, TRADITIONAL MEDICINES & HEALTH SUPPLEMENTS CONSUMED AT THE SAME TIME AND/ OR IN THE LAST 3 MONTHS BEFORE THE ADR) (Additional pages may be attached)						
Product name, active ingredient & strength	Dosage form	Dosage regimen (dosage, frequency & route)	Product details: Batch No. (if known)	Date started	Date stopped	Prescribed for
1.						
2.						

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3.			
4.			
5.			

(5) OTHER RELEVANT INFORMATION (Additional pages may be attached)

Relevant Medical History (Include Status, Smoking, Renal/ Hepati (For congenital abnormalities, please st during pregnancy and the last n	Relevant Investigations (Rechallenge If Performed/ Laboratory Data)	
(6) REPORTER DETAILS *		
Name:		Signature:
Date:	_ Tel No:	Email:

Thank you for taking the time to complete this form.

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GUIDANCE ON ADR REPORTING

WHAT TO REPORT?

An adverse drug reaction is a response to a drug that is noxious (harmful) and unintended, and which occurs at doses normally used in man for the prophylaxis, diagnosis, or therapy of disease, or for the modification of physiological function.

The Ministry of Health encourages the reporting of **all** suspected adverse reactions to drugs and medicinal substances (including self-medication, traditional medicines or health supplements). In particular, please report:

- All suspected reactions to established products and new medicines regardless of their nature and severity.
- All serious adverse reactions which include reactions suspected of causing death, danger to life, admission to hospital, prolongation of hospitalisation, birth defects, persistent or significant disability or incapacity and if medically significant.

To:

• All suspected drug interactions

HOW TO SUBMIT THE REPORT?

The Suspected Adverse Drug Reaction Report form can be obtained from the National Adverse Drug Reaction Monitoring Centre and the nearest government pharmacy facility (hospital/ health centre). This form should be filled as completely as possible and returned to the address below or to the nearest government pharmacy facility (hospital/ health centre).

SUBMISSION OF FOLLOW-UP REPORTS

Any follow-up information for an ADR that has already been reported can be sent to us in another form or *via* any other modes of reporting. Please state that it is a follow-up report, indicating the date and reference number of the initial report.

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National Adverse Drug Reaction Monitoring Centre (NADRMC)

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