ADVERSE DRUG REACTION AND PRODUCT QUALITY PROBLEM REPORT FORM

NATIONAL ADVERSE DRUG EVENT MONITORING CENTRE

Medicines Control Council,
The Registrar of Medicines, 
Department of Health

In collaboration with the WHO International Drug Monitoring Programme

PATIENT INFORMATION

Name (or initials): .......................................................... Age: ....................... Weight (kg) : ....................... 
Sex:  M  F  DOB : ...... / ...... / ...... Height (cm) : ....................... 

ADVERSE REACTION/PRODUCT QUALITY PROBLEM

Adverse reaction  and/or Product Quality problem  Date of onset of reaction: ....../....../...... 
Time of onset of reaction: ......h.......min  

Description of reaction or problem (Include relevant tests/lab data, including dates):

1. MEDICINES/VACCINES/DEVICES (include all concomitant medicines)

<table>
<thead>
<tr>
<th>Trade Name &amp; Batch No. (Asterisk Suspected Product)</th>
<th>Daily Dosage</th>
<th>Route</th>
<th>Date Started</th>
<th>Date Stopped</th>
<th>Reasons for use</th>
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ADVERSE REACTION OUTCOME (Check all that apply)

dehydration
hospitalisation
other.............
Event reappeared on rechallenge:
Y  N  Rechallenge not done 
Treatment of (reaction):.................................
...

Recovered:   Y   N 
Sequelae:     Y   N 
Describe Sequelae:.................................

required intervention to prevent permanent impairment/damage
...

COMMENTS: (e.g. Relevant history, Allergies, Previous exposure, Baseline test results/lab data)

2. PRODUCT QUALITY PROBLEM:

<table>
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<tr>
<th>Trade Name</th>
<th>Batch No</th>
<th>Registration No</th>
<th>Dosage form &amp; strength</th>
<th>Expiry Date</th>
<th>Size/Type of container</th>
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Product available for evaluation?:  Y  N

REPORTING DOCTOR/PHARMACIST Etc:

NAME: ..........................  QUALIFICATIONS: ..........................
ADDRESS: ..........................  
Date: ..........................
TEL: (......)....................

This report does not constitute an admission that medical personnel or the product caused or contributed to the event.
ADVICE ABOUT VOLUNTARY REPORTING

Report adverse experiences with:
- medications (drugs, vaccines and biologicals)
- medical devices (including in-vitro diagnostics)
- traditional and herbal remedies
- For Adverse Events Following Immunisation (AEFI), please follow the reporting procedure recommended by the Expanded Programme in Immunisation (EPI)

Please report:
- adverse drug reactions to recently marketed products
- serious reactions and interactions with all products
- adverse drug reactions which are not clearly reflected in the package insert.

Report even if:
- you’re not certain the product caused the event
- you don’t have all the details

Report Product Quality Problems such as:
- suspected contamination
- questionable stability
- defective components
- poor packaging or labelling
- therapeutic failures

Important numbers:
Investigational Products and Product Quality Problems:
- (012) 326-4344 to fax a report
- (012) 312-0000 to report by phone
Registered Medicines and Traditional and Herbal remedies:
- (021) 448-6181 to fax a report
- (021) 447-1618 to report by phone
Adverse Events Following Immunisation:
- (012) 312 0110 to phone for information
- (012) 321 9882 to fax a report

Confidentiality: Identities of the reporter and patient will remain strictly confidential.

Your support of the Medicine Control Council’s adverse drug reaction monitoring programme is much appreciated. Information supplied by you will contribute to the improvement of drug safety and therapy in South Africa.

PLEASE USE ADDRESS PROVIDED BELOW-JUST FOLD IN THIRDS, TAPE and MAIL

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DEPARTEMENT VAN GESONDHEID
REGISTRAR OF MEDICINES
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