

**PATIENT'S PARTICULARS**

\*Patient's Name or Initials \_\_\_\_\_ \* Sex:  Male  Female Weight \_\_\_\_\_ Kg Height (cm) \_\_\_\_\_  
 Address or Contact Number: \_\_\_\_\_ \*Age \_\_\_\_\_ Date of Birth (mm/dd/yr) \_\_\_\_\_  
 Medical History/Admitting Diagnosis: \_\_\_\_\_ Ethnic group:  Filipino  Chinese  Caucasian  
 Any Known Allergy:  No  Yes, Specify: \_\_\_\_\_ Pregnancy Status: \_\_\_ No  
 Hospital/facility, if admitted: \_\_\_\_\_ Yes (1<sup>st</sup>, 2<sup>nd</sup>, 3<sup>rd</sup> trimester)

**\*DETAILS OF THE ADVERSE REACTION**

Date of onset: \_\_\_\_\_; \_\_\_\_\_ am, \_\_\_\_\_ pm Do you consider the reaction to be serious?  Yes, if yes indicate why:  No  
 Describe the reaction, including pertinent laboratory data:  
 Patient died due to reaction  
 Involved or prolonged in-patient hospitalization  
 Life threatening  
 Involved persistent or significant disability  
 Congenital anomaly in the newborn  
 Other outcome, please give details:  
**Can this be due to Medication Error?**  No  
 Yes, if yes, which type:  
 \_\_\_\_\_ Prescribing  
 \_\_\_\_\_ Transcription  
 \_\_\_\_\_ Dispensing  
 \_\_\_\_\_ Administration

**Can the adverse reaction be due to :**  
 1. **Product quality defect** \_\_\_ No \_\_\_ Yes, Specify, encircle: color change ; caking; powdering ; counterfeit; odor change; defective container; contaminants; separation of components; undissolved suspension/powder  
 2. **Therapeutic failure:** \_\_\_ No \_\_\_ Yes, Specify, encircle: antimicrobial resistance, drug interaction, poor compliance, counterfeit, expired; improper storage; under-dosing, inappropriate medication; inappropriate route of administration; excipients/preservatives

*Suspected drug product(s) Indicate brand name	Daily Dose	Route	Date started	Date stopped	Reason (s) for using the product (Indication)	Manufacturer and Batch/Lot #

List all other drug/s taken at the same time and/ or 3 months before. If none, check box.  No Other drug/s taken

Brand name of the drug	Daily Dose	Route	Date started	Date stopped	Reason/s for using the drug	Manufacturer and Batch & Lot No.

**\*MANAGEMENT OF ADVERSE REACTION**

Was treatment given?  No  Yes (If yes, please specify): \_\_\_\_\_  
**Outcome:**  
 Recovered (Date of recovery): \_\_\_\_\_  Unrecovered **Other diseases:** \_\_\_liver \_\_\_renal \_\_\_HPN  
 Fatal (Date of death): \_\_\_\_\_  Unknown \_\_\_Diabetes \_\_\_CVS \_\_\_Endocrine \_\_\_Cancer  
**Sequela/e:** (any permanent complications or injuries as a result of the ADR) **Re-challenge?**  Yes **Result** \_\_\_\_\_  
 Yes (Please specify) \_\_\_\_\_  No  Unknown  No

**\* REPORTER'S PARTICULARS**

\*Printed Name of Reporter: \_\_\_\_\_ \*Contact no: \_\_\_\_\_  
 Signature of reporter: \_\_\_\_\_ Email address: \_\_\_\_\_  
 Date reported (mm/dd/yr): \_\_\_\_\_ \*Profession: \_\_\_MD \_\_\_RPh \_\_\_RN \_\_\_Patient \_\_\_Dentist \_\_\_other  
 \*Facility: \_\_\_Clinic \_\_\_Trial site \_\_\_Other



**National Pharmacovigilance Center  
 "Saving Lives Through Vigilant Reporting"**

Send completed form to: ADR Unit, FDA, Civic Drive, Filinvest Estate, Alabang, Muntinlupa, 1781.  
 Or fax to: (02) 807-85-11, c/o The ADR Unit. Send sample, if any, of suspect drug for analysis.  
 Website: www.fda.gov.ph

