"PHARMWATCH" MINISTRY OF HEALTH DRUG MONITORING FORM

HEALTH CARE PROFESSIONALS

ENSURE

SAFER PHARMACEUTICALS

PARTICIPATE IN THE DRUG MONITORING PROGRAMME

Report drug failure and adverse reactions with medications and suspected counterfeit product

An adverse reaction occurs when the patient outcome is:

Death, life-threatening (real risk of dying), hospitalization (initial or prolonged), disability (significant, persistent or permanent), congenital defect, permanent impairment, allergic reactions, gastrointestinal distress.

Report even if:

- You're not certain whether the product caused the adverse reaction
- You don't have all the details

Who can report?

Any health care professional (Physician, Pharmacist, Dentist, Nurse) Any patient who has experienced an adverse drug reaction

Where to report:

After completing, please return this form to: Mr. Andre Dennis

Standards & Regulation Division 9th Floor Ministry of Health 2-4 King Street Tel: 948-4106: fax: 967-1629mail Email: dennisa@moh.gov.jm

or Mrs. Princess Thomas Osbourne Standards & Regulation Division 9th Floor Ministry of Health 2-4 King Street Tel: 948-4106: fax: 967-1629 Email: osbournep@moh.gov.jm or Dr. Maxine Gossell-Williams, Department of Basic Medical Sciences Pharmacology Section, University of the West Indies Tel: 927-2216; fax 977-3823 Email: maxine.gossell@uwimona.edu.jm

For additional information or for reporting online please visit the Ministry of Health's website at www.moh.gov.jm

What happens when the Form is submitted?

Any information provided in this form will be handled confidentially. The identities of the health care professional, patient or any other person reporting will be held in strict confidence and protected to the fullest extent. All reports will be assessed and causality analysis decided by Ministry of Health in due course. It is the ultimate responsibility of MOH to decide how to act on the information. It is also the responsibility of the Ministry to decide whether the incidences of reports will require further evaluation of drug performance. The Ministry will further provide the relevant pharmaceutical company with a summary of its findings and subsequent decision regarding intervention.

Prepared by: Maxine Gossell-Williams Department of Basic Medical Sciences Pharmacology Section

> Princess Thomas Osbourne Standards & Regulation Division Ministry of Health

"PharmWatch" is a collaborative effort between the Ministry of Health and the Pharmacology Section of the University of the West Indies.

2009 May

"PHARMWATCH" DRUG MONITORING FORM									
A. PATIENT DETAILS									
1.Patient Initials: (First, Last)	3.Gender: $\Box M \Box F$	3. Date (mm/dd/yyy	of Birth: y)	4. Ethnicity		5. Weight:((Kg)	6. Height: (cm)		
B. SUSPECTED D	RUG EVENT			•					
7. Outcomes attributed to use of drug (check all that apply):			8. Describe event or problem				9.Date event started (mm/dd/yyyy)		
 Failure of therapy Disability Hospitalisation 									
Death mm/dd/yyyy							10. Date event ended (mm/dd/yyyy)		
Other (describe)									
11. Describe action taken in response (e.g. drug changed, prolonged-therapy, increased dose)			12. Describe other relevant history including abnormal laboratory test results, days of hospitalization.						
C. DRUG INFORM			-						
13.Name of suspected drug (give specific name on package)			14. Dose & Route		15. Indic	ation	16. Batch number if known		
	on puekage)								
17. Name of other drugs taken (give specific name on package)			18. Dose & Route 19. Indication		ation	20. Batch number if known			
	1 0 /								
D. REPORTING H	HEALTH PROF	ESSIONA	AL INFORMA	TION	1				
21. Profession:					24.Telep	hone:			
22. Name:		25. Fax:							
23. Address:			26. Email						
27. Also reported to:									
Signature		Date (mm/dd/yyyy)							

FOR OFFICIAL USE ONLY	Code No	
Received by:	Action taken:	
Date received:		