

MEDICINAL PRODUCT DEFECT REPORTING FORM

Completed form to be returned to: Inspectorate and Enforcement Division, Medicines Authority, Sir Temi Żammit Buildings, Malta Life Sciences Park, San Ġwann SĠN3000 or by e-mail at inspectorate.adm@gov.mt.

Shaded areas to be completed by Medicines Authority Staff	Date:	Time:		
	Reference: MDR	Initials:		
Please complete sections 1 to 6 providing as much information as possible. 1. Report made by				
Name:		Position/Status:		
Organisation:				
Address:				
Telephone No: Work:	Fax:	Home:		
E-mail address:				
2. Product details Product name:				
Supplier (from label):				
Manufacturing Site:				
Marketing Authorisation No:				
Legal status	POM / OTC			
Dosage form:				
Strength:				
Container type/size:				
Batch/Lot No:				
Expiry date (if known):				
First distributed (if known):				
Is sample available for Medicines Authority arranged testing? YES/NO				
3. Reported defect and details of any associated clinical incident.				
Do you consider the suspected defect to be: MINOR / SERIOUS / LIFE THREATENING / DON'T KNOW				



4. Contact that can give further information of any clinical incident.				
Name:		Position/Status:		
Organisation:				
Address				
Telephone No: Work:		Fax:		
E-mail address:				
5 Has manufacturer/supplier been informed?		YES/NO		
6. Other action taken by reporter:				
7. Company Contact				
Name:		Position/Status:		
Company:				
Address				
Telephone No: Work	Ext	Fax:		
e-mail address:				
8. The following details should be obtained/confirmed with the licence holder				
Site of manufacture				
Date of distribution				
Batch size				
Distribution (including other countries)				
Other similar defects				
Retained sample to be tested / examined.				
Name of QP(s) responsible for batch release				



9. Comments o	f Duty Medicines Insp	ector:	
Initials:	Date:	Time:	
10. Comments	10. Comments of Duty Medical Assessor (where applicable)		

11. The following details should completed when available			
Cross ref. to other file(s)	Ref no:		
Defect confirmed?	Y/N		
Recall required?	Y/N		
Drug Alert to be issued?	Y/N		
12. Drug Alert/Recall Details Class	1/2/3/4		
Date			
Reference Number	MDR		
Level	Wholesaler /H	Iospital Pharmacy/Community Pharmacy / Patient	
Distribution (In addition to miscellaneous list)	Hospital Only	/ Hospitals & Pharmacies	
Rapid Alert issued	Y/N		
13. Company Reports			
Initial report received	Y/N	Date:	
Interim report received (if required)	Y/N	Date:	
Closing report received	Y/N	Date:	
14. Administrative details Communication to Competent Authority in Country of Manufacture	Date		
File opened	Date:		
Acknowledgement sent to reporter	Date:		
Closing letter sent to:	Reporter	Date	
	Company	Date	



QD002/08 Appendix 01 Version 01

File closed	Date
Database updated	Date
15. Additional notes	