

MINISTRY OF HEALTH OF THE REPUBLIC OF ARMENIA

Scientific Centre of Drug and Medical Technology Expertise after academic E. Gabrielyan REPORT

OF ADVERSE REACTION OR MANUFACTURING PROBLEM OF MEDICINAL PRODUCT

À. INFORMATION AB	OUT PATIEN	T									
1. Name, surname		2. Age or date of birt			th 3. Sex ☐ Ì ☐ F		4	4. Weight	(KGs)	5. Height(cm)	
B. ADVERSE REACTION OR MANUFACTURING PROBLEM											
				2. Date of event (day/mo/yr)				3. Date of report (day/mo/yr)			
4. Description of adverse reaction	or manufacturing pro	blem					<u> </u>				
5.Adverse reaction diagnosis methods used											
6. Short description and peculiarities of disease, laboratory data					□ smoking □ allergy □ alcohol use □ concomitant diseasec			□ pregnancy□ genetic factors□ organ and system dysfunction			
								□ other			
7. Outcome of adverse reaction											
 □ Recovery without sequela □ Recovery with sequela □ Not recover yet □ Death related to ADR 		□ Disabi □ Requi		apacitention	ty of medical pe or prolonged	ersonnel		ome is ur		irth defect	
C. SUSPECTED MEDI	CINAL PROD	UCT	(S)								
Name, manufacturer, batch number , expiry date	2. Dosage form	3. Dose 4. R			te of 5. Indications for us istration		ons for use		6. Thera from /	py dates (duration) ' to	
D. ASSOSIATED TRE	ATMENT (ex	clude	those	e us	sed to tre	at adve	rse read	ction)			
Name, manufacturer, batch number , expiry date	2. Dosage form	3. Dose sngle/d	se 4. Ro		oute of 5. Indicatinistration		ndications for use		6. Therapy dates (duration) from / to		
E. INFORMATION AB	OUT REPOR	ΓER						1			
1. Name, address, phone				2. Profession				3. Occupation			
				□ Doctor							
				☐ Pharmacist							
				□ Other							

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F. ADDITIONAL INFORM	IATION									
Multiplication of the same medicinal product was administered before? If yes, what kind of adverse reactions were observed?										
,										
□ not administ	arad bafara	□ camo	reactions	□ unknown						
□ not administered before			Teactions	□ dilkilowii						
□ other										
2. Whether event abated after us	e stonned?									
2. Whomor event abased after de	о оторром.									
		□ yes	□ no	☐ unknown						
2 \\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \										
3. Whether event reappeared aft	er reintroduction?									
				The state of the s						
	□ yes	□ no	□ unknown	☐ there was no reintroduction						
4. Whether event abated after do	se reduced?									
	□ yes	□ no	□ unknown	☐ there was no dose reduction						
	□ yes	□ 110	L UTKHOWH	☐ there was no dose reduction						
G. ANALYSIS OF REPOR	RT									
Relation between adverse reaction and suspected medicinal product										
☐ definite ☐ probable	☐ possible ☐ dou	ıbtful □ c	onditional impos	ssible to classify						
·	·			·						
Adverse reaction type										
,			_							
	□ seriou	S	□ expecte	d						
	□ not se	erious	□ unexpec	ted						
			·							
Status of medicinal product										
2. Claras of Modelina product										
☐ registered	☐ not registered	⊔ hur	manitarian aid	□ clinical trial						

Reports can be send to Scientific Center of Drug and Medical Technology Expertise department of monitoring of side and adverse effect of medicinal products by the following address:

0051, Yerevan, 49/5 Komitas av.

Tel: (+374 60) 83-00-73

Hot line: (+374 10) 20 05 05, (+374 96) 22 05 05

e-mail: vigilance@pharm.am

Online version of this report can be found at:

web: http://www.pharm.am

You can help million of patients by reporting about suspected side and adverse effects of medicinal products.