Please complete as many details as possible and forward within one business day

Program Details												
Form Completed By Print Name: ELENA SILKA					Name of Program: TREATMENT OF HCV PATIENTS WITH DONATED "SOVALDI" MEDICINE IN THE REPUBLIC OF							
Print Name:	ARMENIA Name of Organisation:											
Signature:	NATIONAL CENTRE FOR AIDS PREVENTION											
Telephone Num	Date aware of Safety Information: 22/06/2017											
Fax No/Email:	Country of Occurrence of Safety Information ARMENIA											
Patient Details												
DOB: 24/05/1976 (or year of birth): Sex:					☑ Female □ Initials: V.Th. Age:					41		
Drug Details (Provide additional drugs on a separate page)												
Lot/Batch No	Reason For Taking	Stop Date (or (DD/MON/			Start Date (DD/MON/YYYY)	Route		Dose	Drug		Name	
L000726 TZDPD	НЕР С	On-going		09	/06/2017	PO		400MG		SOVALDI		
	НЕР С	On-going		09	/06/2017	РО		60MG		DAKLATASVIR		
Safety Information Details: Please provide a short summary of the adverse event(s) (AE) or other safety information (e.g. reports such as pregnancy, death, hospitalization, overdose, misuse, abuse, medication error, lack of effect, off-label use, occupational exposure. AEs associated with product complaints or AEs in an infant following exposure from breastfeeding). Please include the start and stop dates and the outcome of the event(s) or confirm if the event(s) is/are still ongoing. Please also provide any treatment given to treat the event(s), any relevant medical history and for reports of death include the date of death – continue on another page if necessary. Patient started treatment with sovaldi and daklatasvir on 09/06/2017. At the same day experienced dizziness and palpitation for about 30 minutes. The AEs expired without any medical intervention. Medicines were not stopped. Patient also is on the HIV treatment from 2015. Apart of HCV medicines he was administered also the following HIV medicine: 1. Efavirenz / Emtricitabine / Tenofovir 300/200/600mg												
Has this safety information previously been reported to a Regulatory Authority? Yes □ No ☑ Does the Reporter consider that the event(s) were possibly related to drug? Yes ☑ No □								related to the				
Reporter Details (i.e. who notified you of the above safety information?)												
Is the Reporter a: Doctor ☑ Nurse □ Pharmacist □ Non-healthcare professional (e.g. patient, relative)* □ If the Reporter is a Healthcare Professional (HCP) and they are willing to provide us with their contact information, please record below												
*If the Reporter is a Non-healthcare professional, please confirm if they are willing to provide contact information for their HCP: Yes (Please record HCP details below) No												
HCP Address					HCP Name:							
First Line:					HCP Telephone No/FAX No:							
Town/City:					·							
County/State:					HCP Email:							
Postcode/Zip co	de:											

used to comply with applicable laws and regulations. By personal or sensitive data by MoLHSA in accordance with	providing us with information you are consenting to the control and processing of this the applicable data protection laws and the MoLHSA privacy policy.