

QMS N°: DIS/FOM/129 Rev. N°: 1 Effective date : 28/09/2021 Ref. Doc. : PSM/GDL/011

ADVERSE DRUG REACTION/ADVERSE EVENT FOLLOWING IMMUNIZATIONREPORTING FORM

Type of Report		Serious	Seriousness of ADR/AEFI		Category of Suspected Product			
Initial \Box Follow up \Box	110	Serious	□Not Serious□		Medical pro	oduct	Vaccine	
I.PATIENT INFORM	ATION							
Patient ID/initials:Gender: Male Female Weight(kg)				Patient's Medical History(Provide any relevant medical history and laboratory results including dates (if done):				
Height (m):Pregnancy Status: YES NO Date of birth:/								
Patient Address: Village Cell:								
Sector: Phone N°								
II. INFORMATION ON ADVERSE EVENT(S)								
Brief description of the ADI	R/AEFI:							
(a) Information on Onset:				(d) Adverse Event Evolution/ Outcome:				
Date of ADR/AEFI onset:/(dd/mm/yyyy)				Recovered [Recovered Recovering Recovered with sequelae			
Time of onset: / (hours, Min, Sec) Date ADR/AEFI stopped: / (dd/mm/yyyy)				Not recovered Congenital abnormality Death Unknown				
(b) <u>Severity of the ADR/AEFI</u> : Mild Moderate Severe Unknown				(e)Causality of the ADR/AEFI (If performed):				
Reason for seriousness: Prolonged hospitalization Disability Congenital abnormality Death Life threatening								
(c) Action Taken:				(f) Optional information:				
Drug withdrawn Dose increased Dose reduced Dose not changed Substituted Antidote Other (<i>Specify</i>):				(s) that show □Medication	□Therapeutic Failure (Provide information on medicine (s) or vaccine (s) that showed lack of efficacy □Medication errors (Provide details of medication errors)			
III. INFORMATION	ON SUSPECTED PI	RODUCT						
A. Details of suspected								
Product brand name&manufacturer	Generic name/ /Strength/ Dosage form	Route of Administration	Dose and frequency	Starting Date and Time	Stopping Date and Time	Batch Nº. & Expiry date	Indications (Reason for use)	
Other medicines used at the same time and/ or in the last one month (including herbal medicines)								
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		-		1 /				
B Details of Suspected	Vaccine				Diluent (if an	nlicable)		
B. Details of Suspected Vaccine Date of vaccination		Time of Dose $(1^{st}, 2^{nd},$		Batch/Lot Nº	Diluent (if applicable) Name of diluent Batch/Lot N°& Date & time of re-			
Name of vaccine	Date of vaccination	vaccination	3^{rd} etc.)	&Expiry date	Ivanie of undent	Expiry date	constitution	
77.11	WITCH T	0.04	ALL VA	2108				
IV. REPORTER INFORMATION								
Name of reporter: Qualification:					Phone number			
Health Facility Name:			District:		Report Reference N°			
E mail Address of Reporter:		Contact/Te		Date of report:				
Note: Reporters and paties		strict confident	iality by Rwand				w. Once this form is	
completed please send it to Rwanda FDA via the following email: pv_sm@rwandafda.gov.rw								