This form must be returned to the MAH who provided the product. Please see contact details below:

NOTE: Please use the first the	hree letters of the	e month (e.	.g.: JAN)					
○ New ○ Follow-up					Case No:			
For MAH use only								
Date of receipt:			D	D MM	YYYY			
Received by: (Name and o	rganization – eg C	RO, or comp	pany representative	·)				
Source: O Spontaneous O	Comp. Use OL	it. Othe	er, specify					
Suspect Drug								
Drug, Dosage-form, Strength, Route (Drug, Dosage-form, Strength, Route) (eg. Tab 5mg, oral)	Strength, Route (Drug, Dosage-form, Strength, Route) frequency Batch no. start date: stop dat Dosage-form, Strength, Route)		Therapy stop date: DD/ MM / YYYY	Drug-Event Causal relationship Other, Specify (Causal relationship 1 = Not related, 2 = Related)		use of drug		
			/ /	/ /	<u> </u>			
			/ /	/ /	<u> </u>	<u> </u>		
			/ /	/ /				
			/ /	/ /		<u> </u>		
Action Taken				, ,	<u> </u>			
 None Dose decreased, specify Dose increased, specify Patient Data 		own nently disco grarily interre	ontinued	Not Applicable				
Initials:		Date of E	Rirth:		DD MM YYYY	 7		
Weight:	kg	Height:	, , , , , , , , , , , , , , , , , , ,	cm	Gender O Male			
Adverse Event	3							
Description of Adverse Event	(provide diagnosis	if available)			Event onset date:	DD	MM	YYYY
symptoms and treatment:					Event stop date:	DD	MM	YYYY
				O	utcome of Adverse Eve	nt		
				0	Recovered Recovered with sequelae Not recovered Unknown			
					Death Date of death:	DD	MM	YYYY
					Cause(s) of death:		101101	
Did the event result in hospi	talization or prolon	ged hospitali	zation?	O TES PI	autopsy is performed please ease attach relevant clinical onfirm the event.	forward report. laboratory assess	ments to	

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NOTE: Please use the first three JAN)	letters of the month	(e.g.:				
JAN)				Case No:		
Medical History						
Yes (if yes, please specify) None Unknown						
Other Medication (Medication	on taken during th	e last 3 months	s prior to the	event)		
Drug, Dosage-form, Strength, Route (Drug, Dosage-form, Strength, Route) (eg. Tab 5mg, oral)	Dose & frequency	Therapy Start date: DD / MM /YYYY	Therapy Stop date: DD / MM /YYYY	Indication for use of drug		
		/ /	/ /			
		1 1	/ /			
		/ /	/ /			
		1 1	/ /			
		/ /	/ /			
			/ /			
		/ /	/ /			
		/ /	/ /			
		/ /	/ /			
Has the patient discussed this even healthcare professional?	it with their	○ Yes (if yes,○ No ○ U		e provide their healthcare profess	ional's contact in	nformation below)
Healthcare professional's c	ontact information	1				
Name:			Country:			
Address:			Fax:			
			Phone:			
			Email:			
Reporter						
O Physician O Nurse	○ Pharmacist ○	Patient O R	Relative O	ther, please specify		
Name:			Country:			
Address:			Fax:			
			Phone:			
			Email:			
Pharmacy Name (if applicab	ile)					
Name:			Email:			
Signature						
Sign:			Date of AE	awareness:	DD	MM YYYY

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NOTE : Please	use the	first	three	letters	of the	month	(e.g.:	JAN	١

Case No:	

Drug Safety Data Privacy notice

Your personal data will be processed by the relevant marketing authorisation holder, and its worldwide affiliates, to the extent and for as long as necessary, for the purposes of the compliance with drug safety legal obligations and for storage purposes. If you have any questions related to data privacy, want to contact a Data Protection Officer, or wish to exercise your rights of access, rectification, erasure, and/or restriction (as applicable), please contact the relevant marketing authorisation holder listed below. Learn more about how your personal data is processed, shared, stored, transferred, and retained by accessing the applicable privacy policy on the websites below.

This section applies only if the reporter is the patient or anyone but the prescriber/physician/HCP. Please chose one, as applicable:

- O I grant the MAH permission to contact the prescriber/physician/HCP who treated me/the affected patient when the side effect(s) occurred and authorise him/her to provide data from my medical record related to the event(s) occurred.
- No, I do not grant the MAH permission to contact the prescriber/physician/HCP who treated me/the patient.

If you grant the MAH permission, please provide the information of the prescriber/physician/HCP

2 2 1 1 1 1 1 1						
Contact information						
Name:	Country:					
Address:	Fax:					
	Phone:					
	Email:					