

1. Reporter Details											Initial		□Follow-up	
Reporter Name:								E-mail:						
Contact address:								Telephone number:						
								Fax number:						
Typ	be:							☐ Consumer or other non healthcare professional						
☐ Pharmacist								☐ Other (Specify)						
If reporter is a consumer, have they informed their physician of the								he exposure? ☐ Yes ☐ No						
Has the consumer provided permission to contact their healthcare									professional?)		
If yes, please provide healthcare professional contact details:														
Name: Type:						Telephone:								
Address:				I					Email:	Email:				
2. Patient Details														
Date of birth Age								Height Weight						
(Day/Month/Year)				2.12	50	cm						kg		
				Yrs/mo.										
3. C	3. Company Drug Section													
, , , ,		Strength	Dos	se R	Route		dication Trea		tment	nent Treatment		Lot	Expiry	
									start date		end date			
								(day/mo		nth/year) (day/month/y		year)		
1.														
2.														
3.														
4. D	4. Details of Adverse Event													
	erse Event	ent Start Date Stop		Date nth/year) Hospi			italization		Outcom		e E		vent Causality	
	(day/monui/ye		(day/monu		□ Yes			☐ Recovered / Resol			ved			
			☐ No			nrovida	datas					☐ Not Related☐ Unknown		
				If yes, provide date. hospitalization.			uuies 0	☐ Recovering /Resolving			□ UlikilOWII			
							☐ Not Recovered /Not Resolved ☐ Fatal							
									☐ Fatai					
	reatment	.: 1 - 1 C												
		vided for event: with Company D	rug in resr	onse to	o event	:								
Action taken with Company Drug in response to event:														



6. Concomitant Drugs & Therapies				
7. Medical History				
Patient's concomitant conditions, relevant medical history factors or conditions that may affect the outcome of the pregestational, infections during pregnancy, environmental of	regnancy e.g. alcohol, smoking, other substance con	sumption, hypertension, eclampsia, diabetes including		
8. Completed By				
Name:	Signature:	Date (day/month/year):		