

For use by user-facilities, distributors and manufacturers for

	See OMB statement on reverse
Mfr report #	
UF/Dist report #	
	FDA Use Only

VIEDVVAICH	ATORY reporting		UF/Dist report #		
FDA MEDICAL PRODUCTS REPORTING PROGRAM Page	of			FDA Use Only	
Patient information	C. Suspect medi	cation(s)		THE REAL PROPERTY.	
Patient identifier 2. Age at time of event: 3. Sex 4. Weight	Name (give labeled streng	gth & mfr/labele	r, if known)		
or female lbs Or	#1				
In confidence of birth: male kgs	#2				
B. Adverse event or product problem	2. Dose, frequency & route	used 3.	Therapy date	es (if unknown, give duration)	
. Adverse event and/or Product problem (e.g., defects/malfunctions)	#1	#1			
Outcomes attributed to adverse event (check all that apply)	#2	#2	2		
death congenital anomaly	4. Diagnosis for use (indica	ition)		5. Event abated after use	
required intervention to prevent permanent impairment/damage	#1			stopped or dose reduced #1 Tyes no doesn't	
hospitalization – initial or prolonged other:	#2			арріу	
3. Date of 4. Date of	6. Lot # (if known)	7. Exp. dat	e (if known)	≠2 yes no doesn't	
event this report (mordayryr)	#1	#1		Event reappeared after reintroduction	
Describe event or problem	#2	#2		#1 yes no doesn't	
	9. NDC # - for product proble	ems only (if kno	own)		
		11.		#2 yes no doesn't	
	10. Concomitant medical p	products and th	erapy dates (e.	xclude treatment of event)	
	D. Suspect med	ical dovic			
	1. Brand name	icai uevic	.e		
	2. Type of device				
	3. Manufacturer name & address			4. Operator of device	
				health professional	
				lay user/patient other:	
				Other.	
	6.			5. Expiration date	
	model #				
Relevant tests/laboratory data, including dates	catalog #			7. If implanted, give date	
2. Hoteratt testanaporatory data, including dates				_ (morday/yr)	
	serial #			8. If explanted, give date	
	lot #			- (moday/yr)	
	other #				
	9. Device available for eva	aluation?	(Do not se	and to FDA)	
	yes no	retu	rned to manufa	cturer on	
	10. Concomitant medical	products and t	therapy dates (exclude treatment of event)	
7. Other relevant history, including preexisting medical conditions (e.g., allergies,					
race, pregnancy, smoking and alcohol use, hepatic/renal dysfunction, etc.)					
	E. Initial report	er	THE RESERVE	ALTERNATION OF	
	1. Name & address	pho	one #		
Submission of a report does not constitute an admission that medical personnel user facility	2. Health professional?	3. Occupa	tion	4 Initial reporter also sent report to FDA	



distributor, manufacturer or product caused or contributed to the event.

yes no unk

Medication and Device Experience Report

continued)

Submission of a report does not constitute an admission that medical personnel, user facility, distributor, manufacturer or product caused or contributed to the event.

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U.S. DEPARTMENT OF HEALTH AND HUMAN SERVICES
Public Health Service • Food and Drug Administration

FDA Use Only

Refer to guidelines for specific instructions

or use b	y user facili	ty/distributor-devi	ces only	H. Device r	manufacturers o	only
ck one		2. UF/Dist report nui	nber	Type of reportal	ble event	2. If follow-up, what type?
user facility distributor		death		correction		
User facility or distributor name/address			serious injur	у	additional information	
			malfunction	(see guidelines)	response to FDA request	
				other:		device evaluation
				3. Device evaluate	ed by mfr?	4. Device manufacture date
			not returned to mfr. (mayr)			
Contact person				aluation summary attached	5. Labeled for single use?	
				or provide co	age to explain why not) ode:	yes no
5. Date user facili became aware	ty or distributor	7. Type of report 8.	Date of this report			
(mo/day/yr)	or event	initial	(madayyr)	6. Evaluation code	es (refer to coding manual)	
		follow-up #		method	_	
Approximate age of device		em codes (refer to coding m	nanual)	11.001.00		
ugo or uorroo	patient code	-	-	results	-	
	device			condusions		
	code			4		
11. Report sent to	FDA?	12. Location where event		7. If remedial act	ion initiated	8. Usage of device
yes	no/day/yr)	hospital home	outpatient diagnostic facility	check type		initial use of device
		nursing home	ambulatory surgical facility	recall	notification	reuse
13. Report sent to	manufacturer?	outpatient treatment facility	ou.g.our racinty	repair	inspection	unknown
	no/day/yr)	other:		replace	patient monitoring	If action reported to FDA under
			specify	relabeling	modification/	21 USC 360i(f), list correction/remova
luracturer	name/address	9		other:	adjustment	reporting number:
G. All mar	nufacturers					
		mfring site for devices)	2. Phone number			
			3. Report source			
			(check all that apply			
			foreign study			
			literature			I De la Company
			consumer			
			health			
4. Date received to	by manufacturer	5.	professional			
(morday/yr)		(A)NDA #	user facility			
6. If IND, protoco	ol #	INO #	company representative			
		PLA #	distributor			
7. Type of report		pre-1938 yes	other:			
(check all that		OTC yes				
5-day	15-day	product				
	periodic	8. Adverse event term(s				
mitial	follow-up #					
9. Mfr. report nu	ımber					