U.S. Department of Health and Human Services



For VOLUNTARY reporting of adverse events, product problems and product use errors

Page	of	

Form Approved: OMB I	No. 0910-0291, Expires: 10/31/0
	See OMB statement on revers

Triage unit sequence #

	Reporting Program	n		Page		of							
. PATIENT INI						D. SUSPEC	T PROD	OUCT(S)					
Patient Identifier	2. Age at Time of Eve Date of Birth:	F	emale Iale	or	lb kg	1. Name, Streng			product label)				
In confidence B. ADVERSE F	I VENT, PRODUCT				Ng	#2							
heck all that apply:	,					2. Dose or A	mount		Frequency		!	Route	
Adverse Even	=	olem (e.g., defects/r n Different Manufac		*		#1 #2							
Outcomes Attribu (Check all that app	ted to Adverse Event					3. Dates of Use	(If unknov	vn, give duration	on) from/to (or	5. Eve r	nt Abated	d After	Use
Death:	(mm/dd/yyyy)	_ Disability or F	Permanent	Damage		best estimate	·) `			Stop #1	yes	Dose Ro	educed? Doesn't
Life-threatening		Congenital A	•			#2				-			Apply Doesn't
	- initial or prolonged			nt Medical Ever	its)	4. Diagnosis or	Reason fo	or Use (Indica	tion)	#2 📙	Yes _	_ No	Apply
	vention to Prevent Perm			-		#1		,	,		nt Reapportroduction		After
Date of Event (mm	n/dd/yyyy)	4. Date of this Re	eport (mn	n/dd/yyyy)		#2				- #1 🗌	Yes	No	Doesn't
Describe Event, P	roblem or Product Use	Error				6. Lot #		7. Expiration	on Date	#2	Yes	No	Doesn't
						#1		#1		9. NDC	# or Uni	ique ID	,
						#2		#2					
						E. SUSPEC		CAL DEVI	CE				
						1. Brand Name							
						2. Common De	vice Name						
						3. Manufacture	r Name, Ci	ty and State					
						4. Model #		Lot #	#		1 —		of Device Professional
						Catalog #		Expi	ration Date (r	nm/dd/yyyy	v)		er/Patient
						Serial #		Othe	er#			Other:	
						6. If Implanted,	Give Date	(mm/dd/yyyy)	7. If Ex	cplanted, G	ive Date	: (mm/c	dd/yyyy)
						8. Is this a Sing	No					Patien	t?
						9. If Yes to Item	No. 8, En	ter Name and	I Address of I	Reprocess	or		
Relevant Tests/La	boratory Data, Includir	ng Dates											
						F. OTHER	(CONCC	MITANT)	MEDICAL	PRODU	CTS		
						Product names	and thera	apy dates (exc	clude treatmer	nt of event)			
Other Relevant His	story, Including Preexi	sting Medical Con	ditions (e.	.g., allergies,		C DEBOR	FED -/0-		ntinliha na	110	المحماد		
race, programer, or	moning and diconor dec,	invertigation of the second	no, oto. _/			G. REPORT 1. Name and Ad		e comiaei	ntianty sec	FIIOII OII	Dack		
						Phone #			E-mail				
DRODUCT_A	VAII ARII ITV					2. Health Profe	ssional?	3. Occupation	 n		4. Also	Repor	ted to:
O. PRODUCT A roduct Available fo	r Evaluation? (Do not s	end product to FDA)			Yes	No				l	/anufac	
Yes No		lanufacturer on:	-			5. If you do NO	T want you	ur identity dis	closed		_	Jser Fac	•
			(mm	/dd/vvvv)	_	to the manuf	acturer, pl	ace an "X" in	this box:		∐ D	istribut	or/Importer

(mm/dd/yyyy)

ADVICE ABOUT VOLUNTARY REPORTING

Detailed instructions available at: http://www.fda.gov/medwatch/report/consumer/instruct.htm

Report adverse events, product problems or product use errors with:

- Medications (drugs or biologics)
- Medical devices (including in-vitro diagnostics)
- Combination products (medication & medical devices)
- Human cells, tissues, and cellular and tissue-based products
- Special nutritional products (dietary supplements, medical foods, infant formulas)
- Cosmetics

Report product problems - quality, performance or safety concerns such as:

- Suspected counterfeit product
- Suspected contamination
- · Questionable stability
- · Defective components
- · Poor packaging or labeling
- Therapeutic failures (product didn't work)

Report SERIOUS adverse events. An event is serious when the patient outcome is:

Death

-Fold Here-

- · Life-threatening
- · Hospitalization initial or prolonged
- · Disability or permanent damage
- · Congenital anomaly/birth defect
- Required intervention to prevent permanent impairment or damage
- Other serious (important medical events)

Report even if:

- · You're not certain the product caused the event
- · You don't have all the details

How to report:

- Just fill in the sections that apply to your report
- Use section D for all products except medical devices
- · Attach additional pages if needed
- Use a separate form for each patient
- Report either to FDA or the manufacturer (or both)

Other methods of reporting:

- 1-800-FDA-0178 -- To FAX report
- 1-800-FDA-1088 -- To report by phone
- www.fda.gov/medwatch/report.htm -- To report online

If your report involves a serious adverse event with a device and it occurred in a facility outside a doctor's office, that facility may be legally required to report to FDA and/or the manufacturer. Please notify the person in that facility who would handle such reporting.

-Fold Here-

If your report involves a serious adverse event with a vaccine call 1-800-822-7967 to report.

Confidentiality: The patient's identity is held in strict confidence by FDA and protected to the fullest extent of the law. FDA will not disclose the reporter's identity in response to a request from the public, pursuant to the Freedom of Information Act. The reporter's identity, including the identity of a self-reporter, may be shared with the manufacturer unless requested otherwise.

The public reporting burden for this collection of information has been estimated to average 36 minutes per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to:

Department of Health and Human Services Food and Drug Administration - MedWatch 10903 New Hampshire Avenue Building 22, Mail Stop 4447 Silver Spring, MD 20993-0002 Please DO NOT RETURN this form to this address. OMB statement:
"An agency may not of

"An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control number."

U.S. DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

FORM FDA 3500 (10/05) (Back)

Please Use Address Provided Below -- Fold in Thirds, Tape and Mail

DEPARTMENT OF HEALTH & HUMAN SERVICES

Public Health Service Food and Drug Administration Rockville, MD 20857

Official Business

Penalty for Private Use \$300

BUSINESS REPLY MAIL

FIRST CLASS MAIL PERMIT NO. 946 ROCKVILLE MI

MEDWATCH

The FDA Safety Information and Adverse Event Reporting Program Food and Drug Administration 5600 Fishers Lane Rockville, MD 20852-9787



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