FDA USE ONLY

Triage unit sequence #

MEDWATCH

The FDA Safety Information and

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

Form Approved: OMB	No. 0910-0291, Expires: 6/30/2015	
	See PRA statement on reverse.	

Adverse Event	Reporting Program		Page	1 of <u>3</u>						
A. PATIENT IN	NFORMATION			2.	Dose or Amount		Frequency	Route		
1. Patient Identifier	2. Age at Time of Event or	3. Sex	4. Weight	#1						
	Date of Birth:	Female	lb					<u> </u>		
		Male	or kg	#2						
In confidence		_	kgkg				L	<u> </u>		
B. ADVERSE EVENT, PRODUCT PROBLEM OR ERROR Check all that apply:			3. Dates of Use (If unknown, give duration) from/to (or best estimate)					5. Event Abated After Use Stopped or Dose Reduced?		
1. Adverse Even		g., defects/malfund	ctions)	#1				#1 🔲	Yes No Does	
	Error Problem with Differ	-	•	#2	, , ,			#2 🗖	Yes No Does	
	uted to Adverse Event			11	agnosis or Reason fo	r Use	(Indication)		Apply Apply	
(Check all that ap		bility or Permanent	Damaga	#1					nt Reappeared After ntroduction?	
	(mm/dd/yyyy)	•	-	#2				_ #1 [_]	Yes No Does	
Life-threatenin	_	genital Anomaly/Bi		6. Lo	t #	7 F	piration Date	⊢ #2 ┌┐	Yes No Does	
_ ·	n - initial or prolonged Othe vention to Prevent Permanent	• •	nt Medical Events)	#1	· ·	#1	phanon bate	9 NDC	Apply C # or Unique ID	
3. Date of Event (m		e of this Report (,	#2		#2		- 0.1100	, , or ornado in	
3. Date of Event (III	4. Da	e or this Report (miniaa/yyyy)	E.	SUSPECT MEDIC	CALI	DEVICE			
5. Describe Event,	Problem or Product Use Erro	or		1. Br	and Name					
				2. C c	ommon Device Name			2b	. Procode	
				3 Ma	anufacturer Name, Cit	v and	State			
				"		.y				
				4. Mc	odel #	L	ot #		5. Operator of Device	
									Health Profession	
<u> </u>				Ca	italog #	E	xpiration Date (n	nm/dd/vvv\	Lay User/Patient	
							,	3333		
6. Relevant Tests/L	aboratory Data, Including Da	ıtes			rial #	+	nique Identifier (HDN #	Other:	
				"	and w	١	inque iuentinei (ODI, II		
6. Relevant Tests/L						((-)	-(6) 7 IST		G: P-4- / /d-//	
				6. 11	mplanted, Give Date	(mm/a	<i>0/yyyy)</i>	xpianted,	Give Date (mm/dd/yyyy	
				11	this a Single-use Dev	ice tha	at was Reproces	sed and R	teused on a Patient?	
					_ Yes No ✓es to Item No. 8, Enter	Nama	and Address of E			
] 3. 11 1	res to item No. 6, Enter	Name	and Address of F	teprocesso	л	
7. Other Relevant H	listory, Including Preexisting egnancy, smoking and alcohol	Medical Conditions use, liver/kidney p	ons (e.g., roblems, etc.)							
				F. (OTHER (CONCO	ATIN	NT) MEDICA	L PROD	UCTS	
					uct names and thera					
				G	REPORTER (See	confi	dontiality soc	tion on	hack)	
					me and Address	com	acmanty sco	non on .	Jacky	
C. PRODUCT	AVAILABILITY for Evaluation? (Do not send.)			Na	ıme:					
l	`	,		Ad	ldress:					
Yes No	Returned to Manufacture	er on:(mn	n/dd/yyyy)				_			
D. SUSPECT P				Phor			Si E-mai		ZIP:	
1. Name, Strength, #1 Name:	Manufacturer (from product la	bel)		[π		E-mai	•		
Strength:										
Manufacturer:	17'			Ш	alth Professional? 3	. Occı	ıpation		4. Also Reported to:	
#2 Name:					Yes No					
Strength: Manufacturer:					you do NOT want your the manufacturer, plac			¬	Distributor/Import	
1				11			[

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
- Food (including beverages and ingredients added to foods)

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 (devices)
- · 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 -Fold Herewho would handle such reporting.

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 information in this box applies only to requirements of the Paperwork Reduction Act of 1995

The burden time for this collection of information has been estimated to average 36 minutes per response, including the time to review instructions, search existing data sources, gather and maintain the data needed, and complete and review 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 Office of Chief Information Officer Paperwork Reduction Act (PRA) Staff PRAStaff@fda.hhs.gov

Please DO NOT RETURN this form to the PRA Staff e-mail to the left.

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

FORM FDA 3500 (2/13) (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

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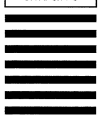
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MEDWATCH

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





U.S. Department of Health and Human Services

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The FDA Safety Information and

(CONTINUATION PAGE) For VOLUNTARY reporting of

adverse events and product problems

Page 3 of 3 **Adverse Event Reporting Program** B.5. Describe Event or Problem (continued) B.6. Relevant Tests/Laboratory Data, Including Dates (continued) B.7. Other Relevant History, Including Preexisting Medical Conditions (e.g., allergies, race, pregnancy, smoking and alcohol use, hepatic/renal dysfunction, etc.) (continued) F. Concomitant Medical Products and Therapy Dates (Exclude treatment of event) (continued)