

DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

MEDWATCH

The FDA Safety Information and **Adverse Event Reporting Program**

Form FDA 3500

Form Approved: OMB No. 0910-0291, Expires: 06-30-2025 See PRA statement on page 5.

	FDA USE ONLY
Triage unit sequence #	
FDA Rec. Date	

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

Note: For date prompts of "dd-mmm-yyyy" please use 2-digit day, 3-letter month abbreviation, and 4-digit year; for example, 01-Jan-1900.

A. PATIENT INFORMATION
1. Patient Identifier (In confidence) 2. Age
B. ADVERSE EVENT, PRODUCT PROBLEM
1. Type of Report (check all that apply) Adverse Event Product Use/Medication Error Product Problem (e.g., defects/malfunctions) Problem with Different Manufacturer of Same Medicine Other Serious or Important Medical Events Congenital Anomaly/Birth Defects
3. Date of Event (e.g., 01-Jan-1900) 4. Date of this Report (e.g., 01-Jan-1900)
5. Describe Event, Problem or Product Use/Medication Error Characters Remaining (max. 4,000): (field continues on next page)

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

^{*} Please see instructions

6. Relevant Test/Laboratory Data	Date (e.g., 01-Jan-1900)	Relevant	Test/Laboratory Data	Date (e.g., 01-Jan-1900)
Additional comments			Character	rs Remaining (max. 2,000):
				5(, ,
7 Other Belgrent History Including Bure	wisting Madical Condition	/1		
7. Other Relevant History, Including Pree problems, etc.)	existing Medical Conditio	ns (e.g., al	ergies, pregnancy, tobac Character	rs Remaining (max. 2,000):

	C. PR	ODUCT	AVAILABILIT	Y		
1. Product Available for Evaluation? (Do not send product Yes No Returned to Manufacturer on (e.g., 0					Do you have a picture of the product? (Check if you are including a picture) Yes	
D. SUSPECT PRODUCTS						
SUSPECT PRODUCT #1						
This report involves: Cosmetic	Dietary supplem	nent	Food/medic	al food	Other	
1. Name, Strength, Manufacturer/Con).			
Product Name		Stren	gth	Unit		
NDC # or Unique ID	Manufacturer/ Comp	mpounder Name			Lot #	
2. Dose or Amount		Frequency Route				
Unit		Other Frequency			Other Route	
3. Treament Dates/Therapy Dates (given by the state of th			· · · · · · · · · · · · · · · · · · ·	top) or da	te of dose reduction.)	
Therapy started on Therapy stopped (e.g., 01-Jan-1900) (e.g., 01-Jan-190			Duration	Unit		
(e.g., 01-3an-1900) (e.g., 01-3an-190	(e.g., 01-3an-19	00)				
Is therapy still on-going? Yes						
4. Diagnosis for use (indication)		5 Prod	uct Type (chec	k all that a	pply) 6. Expiration Date (e.g., 01-Jan-190	
- Elaginosis for acc (malcauch)			TC	Generic		
			ompounded	iar		
7. Event Abated after use Stopped or	Dose Reduced?		it Reappeared a			
Yes No Doesn't apply			es No	Doesn'		
		Ь.			. 455.9	
SUSPECT PRODUCT #2 This report involves: Cosmetic	Dietary supplem	ont	Food/medic	al food	Other	
1. Name, Strength, Manufacturer/Con				ai ioou	Outer	
Product Name	ipoundor (nom prod	Stren		Unit		
		7 S. 18 S.				
NDC # or Unique ID	Manufacturer/ Comp	oounder	ounder Name Lot #			
2. Dose or Amount		Frequency			Route	
2. Dose of Amount		Freque	ПСУ		Route	
Unit		Other Frequency			Other Route	
			Toquolioy			
3. Treament Dates/Therapy Dates (give best estimate of length of treatment (start/stop) or date of dose reduction.)						
Therapy started on Therapy stopped on Dose reduced on OR Duration Unit						
(e.g., 01-Jan-1900) (e.g., 01-Jan-190	00) (e.g., 01-Jan-19	00)				
Is therapy still on-going? Yes No						
4. Diagnosis for use (indication) 5. Product Type (check all that apply) 6. Expiration Date (e.g., 01-Jan-19						
		OTC Generic Compounded Biosimilar				
7. Event Abated after use Stopped or Dose Reduced?		8. Event Reappeared after Reintroduction?				
Yes No Doesn't apply			t apply			

E. SUSPECT MEDICAL DEVICE					
1. Brand Name		2a. Common Device Name	2b. Procode		
3. Manufacurer Name, City an	d State	1			
	T	12			
4. Model #	Lot #	Catalog #	Expiration Date (e.g., 01-Jan-1900)		
Serial # U	nique Identifier (UDI) # 5. (Operator of device			
	[•	Consumer Other		
6a. If Implanted, Give Date (e.	g., 01-Jan-1900) 6b. If Explar	nted, Give Date (e.g., 01-Jan-1900)			
7a. Is this a single-use device		r Name, Address of Reprocessor	8. Was this deviced ever serviced		
that was reprocessed and reused on a patient?			by a third-party servicer? Yes No Unknown		
Yes No					
	F. OTHER (CONC	MITANT) MEDICAL PRODUCTS			
1. Product names and therapy	dates (Exclude treatment of e	event)			
Product Name	Th	nerapy Start Date (e.g., 01-Jan-1900)	Therapy End Date (e.g., 01-Jan-1900)		
1.					
2.					
3.					
4.					
5.					
6.					
7.					
8.					
9.					
10.					
	G. REPORTER (See	confidentiality section on next pag	e)		
1. Name and Address Last Name		First Name			
Lastivame		I list Name			
Address					
City	Sta	te/Province/Region	ZIP/Postal Code		
Country					
Country					
Phone #	mail				
	. Occupation		4. Also Reported to:		
∐Yes ∐No			Manufacturer/Compounder User Facility		
			Distributor/Importer		
5. If you do NOT want your identity disclosed to the manufacturer, please mark this box:					
-	-				

ADVICE ABOUT VOLUNTARY REPORTING

Detailed instructions available at:

https://www.fda.gov/safety/medwatch-forms-fda-safety-reporting/instructions-completing-form-fda-3500

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

- Medications(drugs or biologics)
- Medical devices (including diabetes glucose-test kit, hearing aids, breast pumps, and many more)
- Combination products(medication & medical devices)
- Blood transfusions, gene therapies, and human cells and tissue transplants (for example, tendons, bone, and corneas)
- Special nutritional products(dietary supplements, medical foods, infant formulas)
- Cosmetics (such as moisturizers, makeup, shampoos and conditioners, face and body washes, deodorants, nail care products, hair dyes and relaxers, and tattoos)
- 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
- 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
- Just fill in the sections that apply to your report How to 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)

How to submit report:

- To report by phone, call toll-free: 1-800-FDA (332)-1088
- To fax report: 1-800-FDA (332)-0178
- To report online: www.fda.gov/medwatch/report.htm

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.

Where to submit adverse events related to the following products:

- If your report involves an animal drug, device, pet food and livestock feed problems, go to http://www.fda.gov/vetproductreporting
- If your report involves a health problem or a product problem with a tobacco product, go to https://www.safetyreporting.hhs.gov or call 1-877-287-1373 to report.
- If your report involves an adverse event with a vaccine, go to http://vaers.hhs.gov to report or call 1-800-822-7967.

Confidentiality:

The patient's identity is held in strict confidence by FDA and protected to the fullest extent of the law. 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 40 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 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 above.

OMB statement:

"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