

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

MEDWATCH FORM 3500A

For use by user-facilities, importers, distributors and manufacturers for MANDATORY reporting

Form Approved: OMB No. 0910-0291

Expires: 6-30-2025

See PRA statement on page 6.

FDA USE ONLY					
Mfr report #					
UF/Importer Report #					
Exemption/Variance #					

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 Identif	ier (In confidence)			2. Age Year(s) Month(s)	Week(s) Day(s)	or Date of Birth (e.g., 01-Jan-1900)			
	patient's sex at birth n person has or was i birth).		SECTION REMOVED						
4. Weight lb kg	5. Ethnicity (Check Hispanic/Lati Not Hispanic	no	6. Race (check all that a American Indian/A Asian Black or African A	Native	Native Hawa Other Pacifi White				
		B. AD	VERSE EVENT OR P	RODUCT PRO	OBLEM				
Type of Report (check all that apply) Adverse Event Product Problem (e.g., defects/malfunctions)			2. Outcome Attributed Death – Date of death	eath <i>(01-JAN-19</i> 0 itial or prolonged mportant	00): Red Per Dis	Required Intervention to Prevent Permanent Impairment/Damage Disability or Permanent Damage Congenital Anomaly/Birth Defects			
3. Date of Event	(01-JAN-1900)	4. Date of	this Report (01-JAN-190	00)					

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

* Please see instructions

5. Describe Event or Problem			
6. Relevant Test/Laboratory Data	Date (01-JAN-1900)	Relevant Test/Laboratory Data	Date (01-JAN-1900)
6. Relevant Test/Laboratory Data	Date (01-JAN-1900)	Relevant Test/Laboratory Data	Date (01-JAN-1900)
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	Date (01-JAN-1900)	Relevant Test/Laboratory Data	Date (01-JAN-1900)
6. Relevant Test/Laboratory Data Additional comments	Date (01-JAN-1900)	Relevant Test/Laboratory Data	Date (01-JAN-1900)
	Date (01-JAN-1900)	Relevant Test/Laboratory Data	Date (01-JAN-1900)
	Date (01-JAN-1900)	Relevant Test/Laboratory Data	Date (01-JAN-1900)
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	Date (01-JAN-1900)	Relevant Test/Laboratory Data	Date (01-JAN-1900)
	Date (01-JAN-1900)	Relevant Test/Laboratory Data	Date (01-JAN-1900)

7. Other Relevant His liver/kidney problems,		isting Medical	Conditions (e.g., al	lergies, pregnancy,	tobacco product use, alcohol use, and
		C. SI	JSPECT PRODU	СТЅ	
SUSPECT PRODUCT					
1. Name, Strength, Ma	anufacturer/Compou	nder	01 11	11.20	
Product Name			Strength	Unit	
NDC # or Unique ID	Mar	ufacturer/Comp	 ounder Name		Lot #
2. List Medical Produc	ct and Treatment Giv	en at the Same	Time of the Event	and Date (Do not i	nclude treatment for initial event)
3. Dose or Amount		Fre	quency		Route
Unit		Oth	er Frequency		Other Route
Omt		Otti	errrequency		Other Route
4. Treament Dates/Th	erapy Dates (give be	st estimate of le	ngth of treatment (st	art/stop) or date of o	dose reduction.)
Therapy started on (e.g., 01-Jan-1900) Therapy stopped on (e.g., 01-Jan-1900) (e.g., 01-Jan-1900)				Duration	Unit
(e.g., 61 can 1555)	(e.g., or our root)	(0.9., 07 047	, 500)		
5. Diagnosis for use ((indication)		6. Product Type (d	heck all that apply)	7. Expiration Date (e.g., 01-Jan-1900)
			отс	Generic	
O Frank Aber 1 5	04	. D. d 10	Compounded		ti2
8. Event Abated after		e Keduced?	9. Event Reappear		
Yes No	Doesn't apply		Yes No	Doesn't apply	

SUSPECT PRODUCT	Γ#2									
1. Name, Strength, Manufacturer/Compounder										
Product Name					Strength		Ut	nit		
NDC # or Unique ID	NDC # or Unique ID Manufacturer/Compou				ınder Name				Lot #	
2. List Medical Produc	ct and Tre	atment Give	en at the Sar	me T	Γime of the Εν	vent	and Date	(Do not i	nclude treatment for ini	tial event)
3. Dose or Amount				-	uency				Route	
Unit			0	ther	Frequency				Other Route	
4. Treament Dates/The Therapy started on	Therapy	stopped on	t estimate of Dose Redu		nth of treatmen		art/stop) or Duration		dose reduction.) Unit	
(e.g., 01-Jan-1900)	(e.g., 01-	Jan-1900)	(e.g., 01-Ja	n-19	900)					
5. Diagnosis for use (indication)			6	6. Product Typ	oe (c	heck all tha	at apply)	7. Expiration Date (e	.g., 01-Jan-1900)
					OTC Generic Compounded Biosimilar					
8. Event Abated after	use Stopp	ed or Dose	Reduced?	9	Event Reap	pear	ed after R	eintrodu	uction?	
Yes No	Doesn'	t apply			Yes	No	Does	n't apply	/	
			D. SUS	SPE	CT MEDICA	4L [DEVICE			
1. Brand Name					2a. Commo	n De	vice Name	9		2b. Procode
3. Manufacurer Name	, City and	State								·
4. Model #		Lot#			Ca	italo	g #			
Expiration Date (01-JA	AN-1900)	Serial #								

Unique Device Identifier (UDI) #						
5. Operator of DeviceHealth Professional Patient/Consumer	6a. If Implanted, C	Sive Date (0	11-JAN-1900)	6b. If Exp	olanted, Give Date (01-JAN-1900)	
Other						
7a. Is this a single-use device that was	7b. If yes, enter the name and address of the reprocessor					
reprocessed and reused on a patient?						
Yes No						
8. Was this device ever serviced	9. Is this Device A	vailable for	r Evaluation?	(Do not se	end to FDA)	
by a third-party servicer?	Yes No		(04 141) 40	00)		
Yes No Unknown			on (01-JAN-19			
10. Concomitant Medical Products and Thera Product Name	py Dates (Exclude)			M 1000)	Therapy End Date (01-JAN-1900)	
1.		тпетару оп	art Date (01-JA	114-1900)	Therapy End Date (07-3AN-1900)	
2.						
3.						
4.						
5.						
6.						
7.						
8.						
9.						
10.						
	E. INITIAL	REPORT	ER		<u>'</u>	
1. Name and Address						
Last Name		First Nam	ne			
Address						
7.441.000						
City	State/Province	Region ZIF	P/Postal Code	Country		
Phone # Email						
			1			
2. Health Professional? 3. Occupation (Selection of Selection)	ct from list)				ent report to FDA	
Yes No			Yes	No	Unknown	

F.	F. FOR USE BY USER FACILITY/IMPORTER (Devices Only)						
1. Check One 2. User Facility/Importer Report Number							
User Facility Importer							
3. User Facility or Importer Name	e/Address	4. Contact Perso	on	5. Phone Number			
		6. Date User Fac	ility or Importer	7. Type of Report			
		Became Awar	e of Event (01-JAN-1900	0)			
8. Date of This Report (01-JAN-19	9. Approximate	Age of Device		-			
10. Adverse Event Problem (Refe	r to coding manual)						
Health Effect – Clinical Code	Health Effect – Impact Co	de Medical De	vice Problem Code	Component Code			
11. Report Sent to FDA?	12. Location Where Ev	ent Occurred	'				
(If Yes, enter date (01-JAN-1900))	Ambulatory Surg	ical Facility Out	tpatient Treatment Facilit	y Other (Specify)			
Yes No	Home	Out	tpatient Diagnostic Facilit	у			
	Hospital	Nui	rsing Home				
13. Report Sent to Manufacturer	2 14. Manufactur	er Name/Address					
(If Yes, enter date (01-JAN-190	00))						
Yes							
No							
	G. AL	L MANUFACTURI	ERS				
1. Contact Office (and Manufactur							
Name	3		Address	Phone Number			
Address							
Compounding Outsourcing Facility	503B? Outsourcing	Facility					
Check box if applicable							
2. Report Source (check all that a	oply)			3. Date Received by			
Foreign Literature He	ealth Professional Co	mpany Representative	е	Manufacturer (01-JAN-1900)			
Study Consumer Us	e Faciltiy Dis	stributor/Importer	Other (Please list,				
4. NDA # ANDA	# IND	#	BLA#	PMA/510(k) #			
	"						
Check all that apply:							
	e-ANDA Pre-1938	OTC Co	ompounded Product				
5. If IND/Pre-ANDA, Give Protoco		Check all that apply)	, p				
,		5-day Periodic	Follow-up #				
	7-day 3	0-day Initial	·				
7. Adverse Event Term(s)			8. Manufacturer Repo	ort Number			
7. Advotos Event form(s)			o. manaradarar respe	Transo.			

H. DEVICE MANUFACTURERS ONLY								
1. Type of Reportable Event (check all that apply.) 2. If Follow-up, What Type?						3. Device Evaluated by Manufacturer?		
Death	Malfund	ction Corr			on	Yes	No	
Serious Injury	Summa	ry Report		Addition	al Information			
	No. of e	vents summ	arized	Respons	se to FDA Request			
Device Evaluation								
4. Device Manufacture	Date (01-	JAN-1900)	5. Labe	eled for Single	Use?	1		
			Y	Yes No				
6. Adverse Event Probl	em (Refe	r to coding m	anual)					
Health Effect – Clinical Code Health E			ct – Impact Code Medical Device Prob		blem Code	Component Code		
Type of Investigation			Investi	igation Findings Inves			nvestigation Conclusions	
7. If Remedial Action In	itiated, C	heck Type				8. Usage o	f Device	
Recall		Relabeling	J	Patient Monitor	ing	Initial	Use of Device	
Repair		Notification	n	Modification/Ad	justment	Reus	se	
Replace Inspection Other: Unknown					own			
If action reported to FDA under 21 USC 360i(g), list correction/ removal reporting number:				10. Related	Report Number			
11. Additional Manufacturer Narrative								

This section applies only to requirements of the Paperwork Reduction Act of 1995. This section applies only to requirements of the Paperwork Reduction Act of 1995. The public reporting burden for this collection of information has been estimated to average 73 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 Office of Chief Information Officer Paperwork Reduction Act (PRA) Staff PRAStaff@fda.hhs.gov

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