

## **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 INFO	RMATION						
1. Patient Identifi	er (In confidence)		7.1.7.1.ENT INT	2. Age		or Date of Birth (e.g., 01-Jan-1900)				
1. I atlent identili	ei (iii coimaence)			Year(s)	Week(s)	Date of Billin (e.g., 01-0an-1900)				
				Month(s)	Day(s)					
0.0				Worth(3)						
	patient's sex at birth person has or was birth).		SECTION REMOVED							
4. Weight	5. Ethnicity (Chec	( one)	6. Race (check all that a	oply)						
lb	Hispanic/Lat		American Indian/A		Native Hawa	aiian/				
	Not Hispanio		Asian		Other Pacifi					
kg			Black or African Ar	nerican	White					
		D 4 D			ODLEM					
4.5. (5.			VERSE EVENT OR P							
	(check all that appl	<b>(</b> )	2. Outcome Attributed to			hat apply)				
Adverse Event			Death – Date of de	am ( <i>ur-Jan-19</i>		and the form of the Bound				
Product Proble			Life-threatening			quired Intervention to Prevent manent Impairment/Damage				
(e.g., defects/n	iaiiuriciioris)		Hospitalization (init		,	ability or Permanent Damage				
			Other Serious or Ir	nportant		=				
			Medical Events		Cor	ngenital Anomaly/Birth Defects				
3. Date of Event (	(01-JAN-1900)	4. Date of	this Report (01-JAN-190	0)						

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)
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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)

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	er i requency		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)	Dose Reduce (e.g., 01-Jan-		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 #2											
1. Name, Strength, Ma	anufacture	er/Compoun	der								
Product Name					Strength		Ut	nit			
NDC # or Unique ID Manufacturer/Compo					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 Compour	nded	Gen Bios	eric imilar			
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. <b>Procode</b>	
3. Manufacurer Name	, City and	State								·	
4. Model #		Lot#			Са	italo	g #				
Expiration Date (01-JA	AN-1900)	Serial #									

Unique Device Identifier (UDI) #									
<ul><li>5. Operator of Device</li><li>Health Professional Patient/Consumer</li></ul>	6a. If Implanted, Give Date (01-JAN-1900) 6b. If Explanted, Give Date (01-JAN-1901)								
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			IN 1000)	Therapy End Date (01-JAN-1900)				
1.		тпетару от	art Date (01-JA	(14-1900)	Therapy End Date (07-3AN-1900)				
2.									
3.									
4.									
5.									
6.									
7.									
8.									
9.									
10.									
	E. INITIAL	REPORT	ΓER		·				
1. Name and Address									
Last Name		First Nan	ne						
Address									
7.441.000									
City	State/Province	/Region ZIF	P/Postal Code	Country					
Phone # Email									
			T						
2. Health Professional? 3. Occupation (Selection of Selection)	ct from list)				ent report to FDA				
Yes No			Yes	No	Unknown				

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 Nar	4. Contact P	erson	5. Phone Number							
						ity or Importer		ype of Report		
				Became A	ware	of Event (01-JAN-1900	9)			
8. Date of This Report (01-JAN-1900) 9. Approximate Age of Device										
10. Adverse Event Problem (Re			-1.01-	NA - di -		See Beekleen Onde	0	and Onda		
Health Effect – Clinical Code	Health	Effect – Impa	ct Code	Medica	ai Devi	ce Problem Code	Compo	onent Code		
	1									
11. Report Sent to FDA?		ocation Whe			0	ations Taratarant Facility	_	Other (0:f-)		
(If Yes, enter date (01-JAN-1900)	'	Ambulatory	Surgica	i Facility	-	atient Treatment Facility		Other (Specify)		
Yes No		Home			-	atient Diagnostic Facilit	у			
		Hospital				ing Home				
13. Report Sent to Manufacture (If Yes, enter date (01-JAN-1		14. Manuf	acturer	Name/Addres	SS					
Yes	900))									
No										
110										
				<b>MANUFACT</b>						
1. Contact Office (and Manufact	uring Site	e for Devices)	or Com							
Name				E	-mail <i>F</i>	Address		Phone Number		
Address										
Address										
Compounding Outsourcing Facili	v 503B2	Outsou	cing Fac	rility						
Check box if applicable	y 303D:	Outsoul	cing i ac	Jiilly						
2. Report Source (check all that	annlu)						2 Da	te Received by		
		ofessional	Comp	any Represen	tative			nufacturer (01-JAN-1900)		
Study Consumer				outor/Importer		Other (Please list)		indiacturer (07-0AN-1900)		
		,								
			I			T				
4. NDA # AND	A #		IND#			BLA#		PMA/510(k) #		
Check all that apply:										
<u>'</u>	re-AND/		1938	OTC		npounded Product				
5. If IND/Pre-ANDA, Give Proto	COI#			e <i>ck all that ap<sub>l</sub></i> av Period		Follow up #				
		5-day	15-da	,	JIC .	Follow-up #				
<b>-</b>		7-day	30-da	ay IIIIIai						
7. Adverse Event Term(s)			8. Manufacturer Repo	ort Nun	nber					

H. DEVICE MANUFACTURERS ONLY										
1. Type of Reportable E	vent (che	ck all that ap	ply.)	2. If Follow-up	o, What Type?	3. Device Eva	luated by Manufacturer?			
Death	Malfunction			Correction	on	Yes	No			
Serious Injury	Summa	ry Report		Additiona	al Information					
	No. of e	vents summ	arized	Respons	e to FDA Request					
				Device E	valuation					
4. Device Manufacture	Date (01-	JAN-1900)	5. Labe	eled for Single	Use?					
			Υ	res No						
6. Adverse Event Probl	em (Refe	r to coding m	anual)							
Health Effect – Clinical Code Health E			ect – Impact Code Medical Device P			blem Code	Component Code			
Type of Investigation			Investigation Findings			Investigation Conclusions				
7. If Remedial Action In	itiated, C	heck Type				8. Usage o	f Device			
Recall		Relabeling		Patient Monitori	ng	Initia	Use of Device			
Repair	Repair Notification Modification/Adjustment						se			
Replace Inspection Other:						Unkn	own			
If action reported to FDA under 21 USC 360i(g) list correction/ removal reporting number:				10. Related						
11. Additional Manufacturer Narrative										

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