UF/Importer Report #

U.S. Department of Health and Human Services Food and Drug Administration	For use by user-facilities, importers, distributors and manufacturers
MedWatch	for MANDATORY reporting
FORM FDA 3500A (10/15)	Page 1 of 3

Page 1 of 3

FORM FDA 3500	JA (10/15)			Page 10	013						FDA Use Only
Note: For date promp			git day, 3-letter	month	11	Dose		Frequenc	y I	Route Used	
abbreviation, and 4-di		e, 01-Jul-2015.			#1						
1. Patient Identifier	0.4		3. Sex	4. Weight	#2						
		r(s)									
	or Date of Birth (e		Female		4. Th	erapy Dates (If u	inknown.	 aive duration	1) from/	9 Event Ab	ated After Use
In Confidence			Male	lb	to	(or best estimate)		-	,		or Dose Reduced?
5.a. Ethnicity (Check		— — — — — — — — — — — — — — — — — — —			#1					#1 🗌 Yes	No Doesn't
single best answer)	· _ ` .	American India		ative	#2		<i>A P P</i> .				apply
Hispanic/Latino		African American	White		5. Dia    #1	agnosis for Use	(Indicatio	on)		#2 🗌 Yes	No Doesn't apply
Not Hispanic/Latir	no	awaiian or Other Pa	acific Islander							10 Event Br	
B. ADVERSE E	VENT OR PRO	DUCT PROBL	EM		#2					Reintrod	eappeared After luction?
1. 🗌 Adverse Ever	nt and/or	Product Problem	(e.g., defects/n	nalfunctions)						#1 🗌 Yes	
2. Outcome Attribute	ed to Adverse Eve	nt (Check all that a	pply)			the Product mpounded?		s the Produc he-Counter?			apply
Death Include date (dd-mmm-yyyy):									No	#2 🗌 Yes	No Doesn't
Life-threatening			or Permanent I	° °	#1						apply
Hospitalization –	1 0		al Anomaly/Birt	h Defects		Ves No	° –				_
Other Serious (Im	ntion to Prevent Per		t/Domogo (Do	vices)	11 '		-	/yyy)   #2	_	_	
3. Date of Event (dd-		4. Date of this R	• •								-
			• •			and Name					
5. Describe Event or					11						
					2. Co	mmon Device N	lame				2b. Procode
					3. Ma	nufacturer Nam	ie, City a	nd State			
					4. Mo	del #		Lot #			Operator of Device
6. Relevant Tests/La	horston, Data Inc	luding Datas	(Continue	on page 3)							Health Professional
0. Relevant Tests/La	iboratory Data, inc	luuling Dates			Ca	talog #		Expiration			] Lay User/Patient
					Se	rial #		Unique Ide			Other
								•	,	·  _	
					6. <b>If I</b>	mplanted, Give	Date (dd-	mmm-yyyy)	7. If Exp	planted, Give I	Date (dd-mmm-yyyy)
				on page 3)		-	-			-	-
7. Other Relevant Hi allergies, pregnand	story, Including Pi sy, smoking and alco					this a single-use processed and r				Yes	No
						es to Item 8, En		•	ss of Re	processor	
			(Continue	on page 3)	10. <b>D</b>	evice Available	for Evalu	uation? (Do	not send t	to FDA)	
C. SUSPECT PI 1. Name, Manufactur		24				Yes 🗌 No	Retu	urned to Man	ufacturer	on:	
#1 – Name and Stren			#1 – NDC # or	Unique ID	11. C	oncomitant Mec	dical Pro	ducts and T	herapy D	ates (Exclude	treatment of event)
#1 – Manufacturer/Cc	ompounder		#1 – Lot #		11						
#2 – Name and Stren	ath		#2 – NDC # or							(Cor	ntinue on page 3)
	gui		#2 - NDC # 01		E	NITIAL REPO	<b>JELEB</b>	)			linue on page 3)
#2 – Manufacturer/Co	ompounder		#2 – Lot #			me and Address					
						Name:			First N	ame:	
2. Concomitant Med	ical Products and	Therapy Dates (E	xclude treatme	nt of event)	Addre	ess:					
					City:			Sta	ate/Provin	ice/Region:	
					Coun	try:		'	ZIP/Po	stal Code:	
			(Continue	on page 3)	Phon	e #:		Ema	ail:		
Submission of a re	eport does not c	onstitute an ad	-		2. Hea	alth ofessional?	3. Occup	bation (Select	from list)		I Reporter Also Sent ort to FDA
personnel, user fa			nufacturer or	r product		Yes No					s 🗌 No 🗌 Unk

caused or contributed to the event.

	A (10/	15) (continue	ed)	Page	2 01 3			
F. FOR USE BY	-			_	H. DEVICE MANU		v	
1. Check One	USER			Report Number	1. Type of Reportable Ev		2. If Follow-up, What Type?	
User Facility Importer				Death				
3. User Facility or Importer Name/Address				Serious Injury		Additional Information		
, ,					Malfunction		Response to FDA Request	
							Device Evaluation	
					3. Device Evaluated by		<ol> <li>Device Manufacture Date (dd-mmm-yyyy)</li> </ol>	
			-		Not Returned to I			
4. Contact Person			5. Phone N	lumber		ation Summary Attached		
6. Date User Facility or 7. Type of Report 8 Importer Became Aware			No (Attach page provide code:	to explain why not) or	5. Labeled for Single Use?			
		ort E	B. Date of This Report (dd-mmm-yyyy)			Yes No		
of Event (dd-mmm-y	ууу)	Initial			6. Event Problem and Ev	valuation Codes (Pofer	to poding manual)	
Follow-up #		·						
9. Approximate	10. <b>Eve</b>	0. Event Problem Codes (Refer to cod		ling manual)	Patient Code	-		
Age of Device	Patient				Device			
	Code		_ <b>-</b>		Code			
	Device Code		-	-	Method			
11. Report Sent to FD			ation Where Ev	/ent Occurred				
enter date (dd-mmn			spital	Outpatient	Results	-		
Yes		Ho	•	Diagnostic Facility	Conclusions			
No			rsing Home	Ambulatory				
<ol> <li>Report Sent to Mai Yes, enter date (dd-</li> </ol>	nufacture	er? (If   📛 👝	tpatient Treatme	Surgical Facility ent	7. If Remedial Action Ini	iated, Check Type	8. Usage of Device	
Yes _		Fac	cility		Recall	Notification	Initial Use of Device	
□ <sup>1</sup>			(Specify)	Repair	Inspection			
14. Manufacturer Name/Address				(opcony)	Replace	Patient Monitoring	Unknown	
					Relabeling	Modification/ Adjustment	9. If action reported to FDA under 21 USC 360i(f), list correction/	
					Other:	Aujustinent	removal reporting number:	
							-	
G. ALL MANUFA	CTUR	ERS			10. Additional Manu	facturer Narrative	and / or 11. Corrected Data	
1. Contact Office (and	Manufac	turing Site for I	Devices)	2. Phone Number				
Name								
Address				3. Report Source (Check all that apply)				
Address				Foreign				
				Study				
Email Address								
			Health Professional					
Compounding Outsourcing Facility 503B? Yes			User Facility					
4. Date Received by 5.								
Manufacturer (dd-m	тт-уууу	) NDA #		Representative				
		_ ANDA #		- Distributor				
6. If IND, Give Protoco	əl #	IND #		- Other:				
		BLA #		-				
7. Type of Report		PMA/ 510(k) #	I					
(Check all that apply)								
5-day 30-da	•	Combina	ation Ves					
		Pro Pre-1						
7-day Peric								
10-day Initia	w/-un #	(	OTC Yes					
10-day Initia		r 8. Adverse	e Event Term(s)	)				
10-day     Initia       15-day     Follo		r 8. Adverse	e Event Term(s)	)				
10-day     Initia       15-day     Follo		er 8. Adverse	e Event Term(s)	)				
10-day Initia     10-day Initia     15-day Isolo	rt Numbe			rk Reduction Act of 1995.	Department of Health and		OMB Statement: "An agency may no	

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:

Please DO NOT RETURN this form to the above PRA Staff email address.

Paperwork Reduction Act (PRA) Staff

PRAStaff@fda.hhs.gov

## MEDWATCH

## FORM FDA 3500A (10/15) (continued)

B.5. Describe Event or Problem (continued)

B.6. Relevant Tests/Laboratory Data, Including Dates (continued)

(CONTINUATION PAGE) For use by user-facilities, importers, distributors, and manufacturers for MANDATORY reporting

Page 3 of 3

Concomitant Medical Products and Therapy Dates (Exclude treatment of event) (For continuation of C.2 and/or D.11; please distinguish)

B.7. Other Relevant History, Including Preexisting Medical Conditions (e.g., allergies, pregnancy, smoking and alcohol use, hepatic/renal dysfunction, etc.) (continued)

Other Remarks