

# MEDWATCH

FORM FDA 3500A (10/15)

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Mfr Report #
UF/Importer Report #
FDA Use Only

Note: For date prompts of "dd-mmm-yyyy" please use 2-digit day, 3-letter month abbreviation, and 4-digit year; for example, 01-Jul-2015.

A. PATIENT INFORMATION			
1. Patient Identifier	2. Age <input type="checkbox"/> Year(s) <input type="checkbox"/> Month(s) <input type="checkbox"/> Week(s) <input type="checkbox"/> Days(s)	3. Sex <input type="checkbox"/> Female <input type="checkbox"/> Male	4. Weight <input type="checkbox"/> lb <input type="checkbox"/> kg
or Date of Birth (e.g., 08 Feb 1925)			
5.a. Ethnicity (Check single best answer) <input type="checkbox"/> Hispanic/Latino <input type="checkbox"/> Not Hispanic/Latino			
5.b. Race (Check all that apply) <input type="checkbox"/> Asian <input type="checkbox"/> American Indian or Alaskan Native <input type="checkbox"/> Black or African American <input type="checkbox"/> White <input type="checkbox"/> Native Hawaiian or Other Pacific Islander			

B. ADVERSE EVENT OR PRODUCT PROBLEM	
1. <input type="checkbox"/> Adverse Event and/or <input type="checkbox"/> Product Problem (e.g., defects/malfunctions)	
2. Outcome Attributed to Adverse Event (Check all that apply)	
<input type="checkbox"/> Death Include date (dd-mmm-yyyy): _____	<input type="checkbox"/> Disability or Permanent Damage
<input type="checkbox"/> Life-threatening	<input type="checkbox"/> Congenital Anomaly/Birth Defects
<input type="checkbox"/> Hospitalization – initial or prolonged	<input type="checkbox"/> Other Serious (Important Medical Events)
<input type="checkbox"/> Required Intervention to Prevent Permanent Impairment/Damage (Devices)	
3. Date of Event (dd-mmm-yyyy)	4. Date of this Report (dd-mmm-yyyy)
5. Describe Event or Problem	
(Continue on page 3)	

6. Relevant Tests/Laboratory Data, Including Dates	
(Continue on page 3)	
7. Other Relevant History, Including Preexisting Medical Conditions (e.g., allergies, pregnancy, smoking and alcohol use, liver/kidney problems, etc.)	
(Continue on page 3)	

C. SUSPECT PRODUCT(S)	
1. Name, Manufacturer/Compounder, Strength	
#1 – Name and Strength	#1 – NDC # or Unique ID
#1 – Manufacturer/Compounder	#1 – Lot #
#2 – Name and Strength	#2 – NDC # or Unique ID
#2 – Manufacturer/Compounder	#2 – Lot #
2. Concomitant Medical Products and Therapy Dates (Exclude treatment of event)	
(Continue on page 3)	

E. INITIAL REPORTER	
1. Name and Address	
Last Name:	First Name:
Address:	
City:	State/Province/Region:
Country:	ZIP/Postal Code:
Phone #:	Email:
2. Health Professional? <input type="checkbox"/> Yes <input type="checkbox"/> No	3. Occupation (Select from list)
4. Initial Reporter Also Sent Report to FDA <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unk	

Submission of a report does not constitute an admission that medical personnel, user facility, importer, distributor, manufacturer or product caused or contributed to the event.

3. Dose	Frequency	Route Used
#1		
#2		
4. Therapy Dates (If unknown, give duration) from/ to (or best estimate) (dd-mmm-yyyy)		9. Event Abated After Use Stopped or Dose Reduced?
#1		#1 <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Doesn't apply
#2		#2 <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Doesn't apply
5. Diagnosis for Use (Indication)		10. Event Reappeared After Reintroduction?
#1		#1 <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Doesn't apply
#2		#2 <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Doesn't apply
6. Is the Product Compounded?	7. Is the Product Over-the-Counter?	
#1 <input type="checkbox"/> Yes <input type="checkbox"/> No	#1 <input type="checkbox"/> Yes <input type="checkbox"/> No	
#2 <input type="checkbox"/> Yes <input type="checkbox"/> No	#2 <input type="checkbox"/> Yes <input type="checkbox"/> No	
8. Expiration Date (dd-mmm-yyyy)		
#1 _____	#2 _____	

D. SUSPECT MEDICAL DEVICE		
1. Brand Name		
2. Common Device Name		2b. Procode
3. Manufacturer Name, City and State		
4. Model #	Lot #	5. Operator of Device <input type="checkbox"/> Health Professional <input type="checkbox"/> Lay User/Patient <input type="checkbox"/> Other
Catalog #	Expiration Date (dd-mmm-yyyy)	
Serial #	Unique Identifier (UDI) #	
6. If Implanted, Give Date (dd-mmm-yyyy)		7. If Explanted, Give Date (dd-mmm-yyyy)
-		-
8. Is this a single-use device that was reprocessed and reused on a patient? <input type="checkbox"/> Yes <input type="checkbox"/> No		
9. If Yes to Item 8, Enter Name and Address of Reprocessor		
10. Device Available for Evaluation? (Do not send to FDA) <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Returned to Manufacturer on: _____		
11. Concomitant Medical Products and Therapy Dates (Exclude treatment of event)		
(Continue on page 3)		

PLEASE TYPE OR USE BLACK INK

### F. FOR USE BY USER FACILITY/IMPORTER (Devices Only)

1. Check One <input type="checkbox"/> User Facility <input type="checkbox"/> Importer		2. UF/Importer Report Number	
3. User Facility or Importer Name/Address			
4. Contact Person		5. Phone Number	
6. Date User Facility or Importer Became Aware of Event (dd-mmm-yyyy)		7. Type of Report <input type="checkbox"/> Initial <input type="checkbox"/> Follow-up # _____	
8. Date of This Report (dd-mmm-yyyy)			
9. Approximate Age of Device		10. Event Problem Codes (Refer to coding manual)	
Patient Code _____ - _____ - _____		Device Code _____ - _____ - _____	
11. Report Sent to FDA? (If Yes, enter date (dd-mmm-yyyy)) <input type="checkbox"/> Yes _____ <input type="checkbox"/> No		12. Location Where Event Occurred <input type="checkbox"/> Hospital <input type="checkbox"/> Outpatient Diagnostic Facility <input type="checkbox"/> Home <input type="checkbox"/> Ambulatory Surgical Facility <input type="checkbox"/> Nursing Home <input type="checkbox"/> Outpatient Treatment Facility <input type="checkbox"/> Other: _____ (Specify)	
13. Report Sent to Manufacturer? (If Yes, enter date (dd-mmm-yyyy)) <input type="checkbox"/> Yes _____ <input type="checkbox"/> No			
14. Manufacturer Name/Address			

### G. ALL MANUFACTURERS

1. Contact Office (and Manufacturing Site for Devices)		2. Phone Number	
Name			
Address		3. Report Source (Check all that apply)	
Email Address		<input type="checkbox"/> Foreign	
Compounding Outsourcing Facility 503B? <input type="checkbox"/> Yes		<input type="checkbox"/> Study	
4. Date Received by Manufacturer (dd-mmm-yyyy)		<input type="checkbox"/> Literature	
6. If IND, Give Protocol #		<input type="checkbox"/> Consumer	
7. Type of Report (Check all that apply)		<input type="checkbox"/> Health Professional	
<input type="checkbox"/> 5-day <input type="checkbox"/> 30-day		<input type="checkbox"/> User Facility	
<input type="checkbox"/> 7-day <input type="checkbox"/> Periodic		<input type="checkbox"/> Company Representative	
<input type="checkbox"/> 10-day <input type="checkbox"/> Initial		<input type="checkbox"/> Distributor	
<input type="checkbox"/> 15-day <input type="checkbox"/> Follow-up # _____		<input type="checkbox"/> Other: _____	
9. Manufacturer Report Number		5. NDA # _____	
		ANDA # _____	
		IND # _____	
		BLA # _____	
		PMA/ 510(k) # _____	
		Combination Product <input type="checkbox"/> Yes	
		Pre-1938 <input type="checkbox"/> Yes	
		OTC <input type="checkbox"/> Yes	
8. Adverse Event Term(s)			

### H. DEVICE MANUFACTURERS ONLY

1. Type of Reportable Event <input type="checkbox"/> Death <input type="checkbox"/> Serious Injury <input type="checkbox"/> Malfunction		2. If Follow-up, What Type? <input type="checkbox"/> Correction <input type="checkbox"/> Additional Information <input type="checkbox"/> Response to FDA Request <input type="checkbox"/> Device Evaluation	
3. Device Evaluated by Manufacturer? <input type="checkbox"/> Not Returned to Manufacturer <input type="checkbox"/> Yes <input type="checkbox"/> Evaluation Summary Attached <input type="checkbox"/> No (Attach page to explain why not) or provide code:		4. Device Manufacture Date (dd-mmm-yyyy) ____ - ____ - ____	
		5. Labeled for Single Use? <input type="checkbox"/> Yes <input type="checkbox"/> No	
6. Event Problem and Evaluation Codes (Refer to coding manual)			
Patient Code _____ - _____ - _____			
Device Code _____ - _____ - _____			
Method _____ - _____ - _____			
Results _____ - _____ - _____			
Conclusions _____ - _____ - _____			
7. If Remedial Action Initiated, Check Type <input type="checkbox"/> Recall <input type="checkbox"/> Notification <input type="checkbox"/> Repair <input type="checkbox"/> Inspection <input type="checkbox"/> Replace <input type="checkbox"/> Patient Monitoring <input type="checkbox"/> Relabeling <input type="checkbox"/> Modification/Adjustment <input type="checkbox"/> Other: _____		8. Usage of Device <input type="checkbox"/> Initial Use of Device <input type="checkbox"/> Reuse <input type="checkbox"/> Unknown	
9. If action reported to FDA under 21 USC 360i(f), list correction/removal reporting number:			

10.  Additional Manufacturer Narrative    and / or    11.  Corrected Data

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(CONTINUATION PAGE)

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

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**MEDWATCH**

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

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, pregnancy, smoking and alcohol use, hepatic/renal dysfunction, etc.) (continued)*

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

Other Remarks