Food and Drug Administration

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

// Ifr Report #			
JF/Importer Report #			

MEDWATCH

PLEASE TYPE OR USE BLACK INK

FORM FDA 350	0A (10/05)			Page	of			FDA Use Only			
A. PATIENT IN	FORMATION				C. SUSPECT PRO	DDUCT(S)		•			
1. Patient Identifier	2. Age at Time		3. Sex	4. Weight	1. Name (Give labeled s		labeler)				
	of Event:		☐ Famala	lbs	#1						
	or ————————————————————————————————————		Female	or	#2						
In confidence	of Birth:		Male	kgs	2. Dose, Frequency & F	Poute Head	3 Thera	upy Dates (If unknown, give duration)			
B. ADVERSE E	VENT OR PRODU	CT PROBLE	М			iodic Oscu	from/t	from/to (or best estimate)			
1. Adverse Even	nt and/or Pro	duct Problem (e	e.g., defects/malf	iunctions)	#1		#1				
	ted to Adverse Event				#2		#2				
(Check all that app	ly)	□ 5 :			4. Diagnosis for Use (In	ndication)		5. Event Abated After Use Stopped or Dose Reduced?			
Death:	(mm/dd/yyyy)		or Permanent Da	Ť	#1			#1 Yes No Doesn't			
Life-threatenir	•		I Anomaly/Birth [#2			Apply Apply			
1 = '	n - initial or prolonged		ious (Important M		6. Lot #	7. Exp. D	ate	#2 Yes No Doesn't			
	rvention to Prevent Perm	· · · · · · · · · · · · · · · · · · ·			#1	#1		8. Event Reappeared After			
3. Date of Event (mr	m/dd/yyyy)	4. Date of This	Report (mm/do	d/yyyy)	#2	#2 #2		Reintroduction? #1 Yes No Doesn't			
5. Describe Event or	Describe Frank or Broklam				9. NDC# or Unique ID			Apply Apply			
5. Describe Event of	Problem							#2 Yes No Doesn't Apply			
					10. Concomitant Medic	al Products a	nd Therapy Date:	s (Exclude treatment of event)			
					D. SUSPECT MEI	DICAL DEV	VICE				
					1. Brand Name		VIOL				
					2. Common Device Name						
					3. Manufacturer Name, City and State						
					5. Manufacturer Name,	Oity and Stat	ic				
					4. Model #	Lo	ot #	5. Operator of Device			
					Catalog #	─────────────────────────────────────					
					Serial #	0	ther #	Other:			
					6. If Implanted, Give Da	ite (mm/dd/yy)	7. If Exp	planted, Give Date (mm/dd/yyyy)			
6. Relevant Tests/Laboratory Data, Including Dates					8. Is this a Single-use Device that was Reprocessed and Reused on a Patient? Yes No						
					9. If Yes to Item No. 8, I	Enter Name a	nd Address of Re	eprocessor			
					10. Device Available for						
				Yes No Returned to Manufacturer on:							
					11. Concomitant Medic	al Products a	nd Therapy Dates	s (Exclude treatment of event)			
7. Other Relevant Hi race, pregnancy, si	story, Including Preexis moking and alcohol use, I	sting Medical Co hepatic/renal dys	onditions (e.g., a function, etc.)	illergies,							
					E. INITIAL REPO	RTER					
					1. Name and Address		Phone #				
						·					
Submission of a	report does not c	onstitute an	admission tl	hat medical	2. Health Professional?	3. Occupat	ion	4. Initial Reporter Also Sent			
personnel, user 1	facility, importer, outed to the event.	distributor, n	nanufacturer	or product	Yes No			Report to FDA Yes No Unk.			

MEDWATCH									r DA GSI	CONLI		
FORM FDA 3500	A (10/05) (continued)		Page _	of	_						
F. FOR USE BY	USER FA	ACILITY/IMPO	ORTER (D			EVICE MANU	FACTURE	RS ONLY	,			
F. FOR USE BY USER FACILITY/IMPORTER (D 1. Check One User Facility Importer 2. UF/Importer R					e of Reportable Ev				Correction		?	
3. User Facility or Impo	orter Name	/Address				Serious Injury Malfunction Other:				Additional Response Device Ev	I Information to FDA R	Request
4. Contact Person 5. Phone Nu				3. Device Evaluated by Manufacturer? Not Returned to Manufacturer Yes Evaluation Summary Attached					ice Manufad 1/yyyy)	cture Date	e	
6. Date User Facility or Importer Became Aware of Event (mm/dd/yyyy)			rt	8. Date of This Report (mm/dd/yyyy)		No (Attach page provide code:	to explain wh	y not) or		eled for Sin	igle Use?	
		Follow-up #			6. Eva	uation Codes (Re	fer to coding	manual)				
9. Approximate Age of Device	10. Event I		des (Refer to coding manual)		Method							
	Patient Code	-	-			Results]-[
	Device Code	-	-			Conclusions]-				
11. Report Sent to FDA	.?	12. Location W	/here Event 0	Occurred	7. If Re	medial Action Init	tiated, Check	Туре	3. Usage o	f Device		
Yes Hospital No				Outpatient Diagnostic Facility Ambulatory Surgical Facility		Recall [Notification Inspection		R	nitial Use of [leuse Inknown	Device	
13. Report Sent to Manufacturer? Yes (mm/dd/yyyy) No Other:			• .		Replace [Relabeling [Patient Mo Modification Adjustment	on/	21 USC	reported to 360i(f), list	correction	der n/	
		Outer.		(Specify)		Other:						
G. ALL MANUFA 1. Contact Office - Nan for Devices)			ring Site	Phone Number 3. Report Source (Check all that apply)	10.	Additional Manu	acturer Nam	auve	and / or	11	Correcte	ed Data
		T-		Foreign Study Literature Consumer Health Professional User Facility								
 Date Received by Manufacturer (mm/d 	d/yyyy)	5. (A)NDA # IND #		Company Representative Distributor								
6. If IND, Give Protocol	I #	STN #		Other:								
7. Type of Report (Check all that apply)		PMA/ 510(k) # Combination										
5-day 30-da 7-day Perior 10-day Initial	dic	Product Pre-1938	Yes Yes									
= ' -	w-up #	OTC Product	Yes									
9. Manufacturer Repor	t Number	8. Adverse Ev	vent Term(s)									

The public reporting burden for this collection of information has been estimated to average 66 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 - MedWatch 10903 New Hampshire Avenue Building 22, Mail Stop 4447 Silver Spring, MD 20993-0002

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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."