

APPENDIX F

Form Approved: OMB No. 0910-0291 Expires: 01/31/81
See OMB statement on reverse

MEDWATCH

THE FDA MEDICAL PRODUCTS REPORTING PROGRAM

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

Page ___ of ___

Mfr report #
UF/Dist report #
FDA Use Only

A. Patient information			
1. Patient Identifier	2. Age at time of event: or Date of birth:	3. Sex <input type="checkbox"/> female <input type="checkbox"/> male	4. Weight ____ lbs or ____ kgs
In confidence			
B. Adverse event or product problem			
1. <input type="checkbox"/> Adverse event and/or <input type="checkbox"/> Product problem (e.g., defects/malfunctions)			
2. Outcomes attributed to adverse event (check all that apply)			
<input type="checkbox"/> death (m/d/yyyy)	<input type="checkbox"/> life-threatening	<input type="checkbox"/> hospitalization - initial or prolonged	<input type="checkbox"/> disability <input type="checkbox"/> congenital anomaly <input type="checkbox"/> required intervention to prevent permanent impairment/damage <input type="checkbox"/> other: _____
3. Date of event (m/d/yyyy)	4. Date of this report (m/d/yyyy)		
5. Describe event or problem			
6. Relevant tests/laboratory data, including dates			
7. Other relevant history, including preexisting medical conditions (e.g., allergies, race, pregnancy, smoking and alcohol use, hepatic/renal dysfunction, etc.)			

C. Suspect medication(s)			
1. Name (give labeled strength & mfr/labeler, if known)			
#1 _____			
#2 _____			
2. Dose, frequency & route used		3. Therapy dates (if unknown, give duration) (m/d/yyyy to best estimate)	
#1 _____		#1 _____	
#2 _____		#2 _____	
4. Diagnosis for use (indication)		5. 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	
6. Lot # (if known)		7. Exp. date (if known)	
#1 _____		#1 _____	
#2 _____		#2 _____	
8. Event reappeared after reintroduction			
#1 <input type="checkbox"/> yes <input type="checkbox"/> no <input type="checkbox"/> doesn't apply			
#2 <input type="checkbox"/> yes <input type="checkbox"/> no <input type="checkbox"/> doesn't apply			
9. NDC # - for product problems only (if known)			
- - - - -			
10. Concomitant medical products and therapy dates (exclude treatment of event)			

G. All manufacturers	
1. Contact office - name/address (& mfring site for devices)	2. Phone number
3. Report source (check all that apply)	
<input type="checkbox"/> foreign	
<input type="checkbox"/> study	
<input type="checkbox"/> literature	
<input type="checkbox"/> consumer	
<input type="checkbox"/> health professional	
<input type="checkbox"/> user facility	
<input type="checkbox"/> company representative	
<input type="checkbox"/> distributor	
<input type="checkbox"/> other: _____	
4. Date received by manufacturer (m/d/yyyy)	5. (A)NDA # _____
6. If IND, protocol #	IND # _____
7. Type of report (check all that apply)	PLA # _____
<input type="checkbox"/> 5-day <input type="checkbox"/> 15-day	pre-1938 <input type="checkbox"/> yes
<input type="checkbox"/> 10-day <input type="checkbox"/> periodic	OTC product <input type="checkbox"/> yes
<input type="checkbox"/> initial <input type="checkbox"/> follow-up # _____	8. Adverse event term(s)
9. Mfr. report number	

E. Initial reporter			
1. Name & address			phone #
2. Health professional? <input type="checkbox"/> yes <input type="checkbox"/> no			
3. Occupation			
4. Initial reporter also sent report to FDA <input type="checkbox"/> yes <input type="checkbox"/> no <input type="checkbox"/> unk			

PLEASE TYPE OR USE BLACK INK



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