APPENDIX F



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

Form Approved:	CMS No. 0919-0291 Expires: 01/31/0 See CMS etylement on revery
Mrreport &	
F/Dist report #	
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THE FOA MEDICAL PRODUCTS REPORT	ING PROGRAM Page	of	L	FDA Use Only
A. Patient information 1. Patient identifier 2. Age at time of event: or Date in confidence B. Adverse event or produ	3. Sex 4. Weight	C. Suspect medicati 1. Name (give labeled strength & r #1 #2 2. Dose, frequency & route used #1 #2 4. Diagnosis for use (indication)	nfr/labeler, if known)	tes (if unknown, give duration) stimula) 5. Event abated after use stopped or dose reduced
life-threatening hospitalization - initial or prolonged 3. Date of event (moldsyyr) 5. Describe event or problem	required intervention to prevent permanent impairment/damage other: 4. Date of this report (modesyly)	#1 #2 6. Lot # (if known) #1 #2 9. NDC # – for product problems or — 10. Concomitant medical product	: : : : : : : : : : :	#1 yes no doesn #2 yes no doesn 8. Event reappeared after reintroduction #1 yes no doesn #1 yes no doesn #2 yes no doesn #2 yes no doesn
		G. All manufacturer 1. Contact office ~ name/address (& mining site for devices	2. Phone number 3. Report source (check all that apply) foreign study iiterature consumer health professional
6. Relevant tests/laboratory data, including	dates	4. Date received by manufacturer imordary(r) 5. If IND, protocol # 7. Type of report (check all that apply) 5-day	272	user facility company representative distributor yes other:
Other relevant history, including preex race, pregnancy, smoking and alcohol use	, hepatic/renal dysfunction, etc.)	E. Initial reporter 1. Name & address	phone #	4 Initial reporter also
admission tha	f a report does not constitute an t medical personnel, user facility,	2. Health professional? 3. O	scupation	4 Initial reporter also sent report to FDA yes no unk

FDA Form 3500A

distributor, manufacturer contributed to the event.