

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

MEDWATCH FORM 3500A

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

Form Approved: OMB No. 0910-0291 Expires: 6-30-2025

See PRA statement on page 6.

FDA USE ONLY					
Mfr report #					
UF/Importer Report #					
Exemption/Variance #					

Note: For date prompts of "dd-mmm-yyyy" please use 2-digit day, 3-letter month abbreviation, and 4-digit year; for example, 01-JAN-1900.

	A. PATIENT INFORMATION
Patient Identifier (In confidence) JD123456	2. Age
3. Sex: Enter the patient's sex at birth (the sex that a person has or was assigned to at birth). Male Female	SECTION REMOVED
4. Weight 160.0(no American Indian/Alaska Native Native Hawaiian/
	B. ADVERSE EVENT OR PRODUCT PROBLEM
1. Type of Report (check all that app. Adverse Event Product Problem (e.g., defects/malfunctions)	□ Death − Date of death (01-JAN-1900): □ Life-threatening □ Required Intervention to Prevent Permanent Impairment/Damage □ Other Serious or Important Medical Events □ Disability or Permanent Damage □ Congenital Anomaly/Birth Defects
3. Date of Event (01-JAN-1900) 17-Jan-2025	4. Date of this Report (01-JAN-1900) 22-Jan-2025

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

* Please see instructions

5. Describe Event or Problem					
On January 19, 2025, the patient experienced acute shortness of breath, dizziness, and chest tightness approximately 30 minutes after taking Drug X for the eighth consecutive day. Symptoms escalated over the next hour, leading to an emergency room visit and subsequent hospital admission. Clinical evaluation confirmed a hypertensive crisis potentially triggered by the medication. No prior history of similar symptoms was reported. After discontinuation of Drug X, the patient's condition stabilized within 48 hours.					
6. Relevant Test/Laboratory Data	Date (01-JAN-1900)	Relevant Test/Laboratory Data	Date (01-JAN-1900)		
6. Relevant Test/Laboratory Data Blood Pressure	Date (01-JAN-1900)	Relevant Test/Laboratory Data	Date (01-JAN-1900)		
	Date (01-JAN-1900)	Relevant Test/Laboratory Data	Date (01-JAN-1900)		
Blood Pressure	Date (01-JAN-1900)	Relevant Test/Laboratory Data	Date (01-JAN-1900)		
Blood Pressure Chest X-ray	Date (01-JAN-1900)	Relevant Test/Laboratory Data	Date (01-JAN-1900)		
Blood Pressure Chest X-ray	Date (01-JAN-1900)	Relevant Test/Laboratory Data	Date (01-JAN-1900)		
Blood Pressure Chest X-ray Basic Metabolic Panel & CBC	Date (01-JAN-1900)	Relevant Test/Laboratory Data	Date (01-JAN-1900)		
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Blood Pressure Chest X-ray Basic Metabolic Panel & CBC	Date (01-JAN-1900)	Relevant Test/Laboratory Data	Date (01-JAN-1900)		

7. Other Relevant History, Includiver/kidney problems, etc.)	ing Preexisting	Medical	Conditions (e.g., a	llergies, pregnan	cy, tobacco product use, alcohol use, and
Drug Y with tobacco					
		C. SI	JSPECT PRODU	CTS	
SUSPECT PRODUCT #1					
1. Name, Strength, Manufacturer	Compounder		To: II		
Product Name Drug X			Strength 10	Unit	S) - GM
NDC # or Unique ID	Manufactu	rer/Comp	ounder Name	GRAIVII	Lot #
TABO II OI OIIIQUO IB	Apex Com		ounder Hame		LOC III
2. List Medical Product and Treat			e Time of the Event	and Date (Do n	ot include treatment for initial event)
					·
Dates of Use: 01/12/2025 - 01/20/20	25Diagnosis for	Use: Hype	ertensionEvent Onset	Date: 01/19/2025	
3. Dose or Amount		Fre	quency		Route
-					
Unit Other Frequency			Other Route		
4. Treament Dates/Therapy Dates	(aive hest estir	nate of le	nath of treatment (st	art/ston) or date	of dose reduction)
Therapy started on Therapy started		e Reduce		Duration	Unit
(e.g., 01-Jan-1900) (e.g., 01-Ja	n-1900) (e.g	., 01-Jan-	1900)		
5 Diamenta famous (indiantias)			C. Door door d. T. co. c. (7 Femination Buts (c. p. 04 Jun 4000)
5. Diagnosis for use (indication)			_		7. Expiration Date (e.g., 01-Jan-1900)
			OTC Compounded	Generic Biosimila	
8. Event Abated after use Stoppe	d or Dose Red	uced?	9. Event Reappea		
Yes No Doesn't apply					

SUSPECT PRODUCT #2						
1. Name, Strength, Manufacture	r/Compounder					
Product Name		Strength	Strength Unit			
NDC # or Unique ID	Manufacturer/C	compounder Nar	oounder Name		Lot#	
2. List Medical Product and Tre	atment Given at the S	Same Time of th	ne Event and	Date (Do not i	include treatment for initial	event)
3. Dose or Amount Frequency				Route		
Unit Othe			ісу		Other Route	
4. Treament Dates/Therapy Date	es (give best estimate	of length of trea	tment (start/sto	op) or date of	dose reduction.)	
	stopped on Dose Re				Unit	
5. Diagnosis for use (indication)		6. Produc	t Type (check	all that apply)	7. Expiration Date (e.g.,	01-Jan-1900)
		OTC	pounded	Generic Biosimilar		
8. Event Abated after use Stopped or Dose Reduced? 9. Event Reappeared after Reintroduction?						
Yes No Doesn't apply						
	D. S	USPECT MEI	DICAL DEV	ICE		
1. Brand Name			2a. Common Device Name			2b. Procode
3. Manufacurer Name, City and	State	ı				I
4. Model #	Lot#	ot # Catalog #				
Expiration Date (01-JAN-1900)	Serial #					

Unique Device Identifier (UDI) #				
5. Operator of Device	6a. If Implanted, C	Give Date (01-JAN-1900)	6b. If Exp	planted, Give Date (01-JAN-1900)
Health Professional Patient/Consumer				
Other	75 16			
7a. Is this a single-use device that was reprocessed and reused on a patient?	7b. If yes, enter the	e name and address of the	e reproces	ssor
Yes No				
8. Was this device ever serviced	0 le this Davice	Available for Evaluation?	(Do not se	and to EDA)
by a third-party servicer?	Yes No		(DO NOT SE	and to I DA)
Yes No Unknown		nanufacturer on (01-JAN-19	00)	
10. Concomitant Medical Products and Thera	py Dates (Exclude	treatment of event)		
Product Name		Therapy Start Date (01-JA	AN-1900)	Therapy End Date (01-JAN-1900)
1.				
2.				
3.				
4.				
5.				
6.				
7.				
8.				
9.				
10.				
	E INITIAL	REPORTER		
1. Name and Address	L. INITIAL	REPORTER		
Last Name		First Name		
Address				
City	State/Province	/Region ZIP/Postal Code	Country	
Phone # Email				
2. Health Professional? 3. Occupation (Selection)	ct from list)			ent report to FDA
Yes No		Yes	No	Unknown
'		1		

F. FOR USE BY USER FACILITY/IMPORTER (Devices Only)						
1. Check One 2. User Facility/Importer Report Number						
User Facility Imp	oorter					
3. User Facility or Imported	er Name/Addres	ss	4. Contact Person			Number
			6. Date User Facili		7. Type of Rep	ort
			Became Aware	of Event (01-JAN-1900)	
8. Date of This Report (01	-JAN-1900)	9. Approximate Age	e of Device			
10. Adverse Event Proble Health Effect – Clinical Cod		<i>ing manual)</i> Effect – Impact Code	Modical Davi	ce Problem Code	Component Code	
Health Ellect - Clinical Coc	ie nealth t	illect – Impact Code	iviedicai Devi	ce Problem Code	component Code	
	10.					
11. Report Sent to FDA? (If Yes, enter date (01-JAN)		cation Where Event Ambulatory Surgica		atient Treatment Facility	Other (Sp	o o ifu)
Yes No	. =	Home		atient Diagnostic Facility		эспу)
☐ fes ☐ No	-] Hospital	= '	ing Home		
			_			
13. Report Sent to Manufa (If Yes, enter date (01-		14. Manufacturer	Name/Address			
Yes	JAIV-1900))					
□No						
				_		
			MANUFACTURE			
1. Contact Office (and Mai	nufacturing Site	for Devices) or Com			- N	
Name			Email A	Address	Phone Num	iber
Address						
Address						
Compounding Outsourcing	Facility 503B2	Outsourcing Fac	cility			
Check box if applicable	racility 303B:	Outsourcing rate	Sility			
	II that apply)				3. Date Receive	d by
2. Report Source (check a		fessional Comp	any Representative			-
Foreign Literature Health Professional Company Representative Manufacturer (01-JAN-1900) Study Consumer Use Facility Distributor/Importer Other (Please list)						(01-3AN-1900)
		y	atol/importor			
4. NDA #	ANDA#	IND#		BLA#	PMA/510(k)#
Check all that apply:	_	_				
Combination product	Pre-ANDA			npounded Product		
5. If IND/Pre-ANDA, Give	Protocol # 6.	Type of Report (Che				
			ay Periodic [Follow-up #		
		7-day 30-da	ay			
7. Adverse Event Term(s)				8. Manufacturer Repo	rt Number	

H. DEVICE MANUFACTURERS ONLY								
1. Type of Reportable Event (cf	eck all that apply.	y.) 2. If Follow-up	o, What Type?	3. Device Eva	aluated by Manufacturer?			
☐ Death ☐ Malfun	Correction	n	Yes	No				
Serious Injury Summ	Additiona	al Information						
No. of	zed Respons	e to FDA Request						
		Device E	valuation					
4. Device Manufacture Date (01	-JAN-1900) 5. L	Labeled for Single	Use?					
		Yes No						
6. Adverse Event Problem (Ref	er to coding manu	ual)	T					
Health Effect – Clinical Code	Health Effect –	- Impact Code	Medical Device Prob	olem Code	Component Code			
Type of Investigation	Inv	vestigation Findings		Investigation	Investigation Conclusions			
7. If Remedial Action Initiated,	Check Type			8. Usage o	of Device			
Recall	Relabeling	Patient Monitori	na		Il Use of Device			
Repair	Notification	Modification/Ad		Reus				
Replace	Inspection	Other:	, 4041110111		Unknown			
9. If action reported to FDA und		<u> </u>	Report Number					
list correction/ removal repo		(g), To. Related	Report Humber					
11. Additional Manufacturer Na	rrative							
This section applies only to requirements of the Paperwork Reduction Act of 1995. This section applies only to requirements of the Paperwork Reduction Act of 1995. 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:								
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Food and Drug Administration								
Office of Chief Information Office								
Paperwork Reduction Act (PRA)	Staff							
PRAStaff@fda.hhs.gov								
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