																	CIO		IS I	FO 	RM
SUSPECT ADVERSE REACTION REPORT																					
303FE	CIADVENSE	LAC	TION KEI	OKI						_							_	_	_	_	
															<u> </u>						
1. PATIENT INITIALS	I decinitary			EACTIO				_	400		TION	ONSI		T							
(first, last)	1a. COUNTRY COSTA RICA	Day	DATE OF BIRTH Month Ye	2a. AG	E 3. SE	EX	3a. WEIGHT 120.00	\vdash	4-6 R ay		onth	-	Year	8-12	Α	PPR	CK ALL ROPRI RSE I	ATE			
PRIVACY			PRIVACY	Year	s Fem	nale	kg			L	Ink			╛			ENT DI		01101	•	
7 + 13 DESCRIBE REACTION(S) (including relevant tests/lab data) Event Verbatim [PREFERRED TERM] (Related symptoms if any separated by commas)									INIVOLVED OR												
Headache [Headache]										INVOLVED OR PROLONGED INPATIENT HOSPITALISATION											
Case Description: ***This is an auto generated narrative***											INVOLVED PERSISTENT OR SIGNIFICANT DISABILITY OR INCAPACITY										
Study ID: 828652-My Healthy Journey										LIFE											
Study description: Trial title: This is a 40 weeks digital patient support program with focus on exercise,										CONGENITAL											
motivation, nutrition & maintaining strategies (only for patients under Liraglutide 3.0 mg).										ANOMALY											
					(0	Contin	ued on Ad	ditio	nal I	nfor	mati	on P	age)	OTHER							
			II. SUSPI	ECT DR	UG(S)) INF	ORMA	TIC	N												
II. SUSPECT DRUG(S) INFORMATION 14. SUSPECT DRUG(S) (include generic name) #1) Saxenda (liraglutide 6 mg/mL) Solution for injection, 6 mg/mL											20. DID REACTION ABATE AFTER STOPPING DRUG?										
15. DAILY DOSE(S) #1) 1.8 mg, qd						B. ROUTE(S) OF ADMINISTRATION 1)Subcutaneous								YES NO NA							
17. INDICATION(S) FOR USE											21. DI R			CTION AR AF							
#1) Obesity (Obesity)																TRO	DUCT	TION	?		
						THERAPY DURATION) Unknown								YES NO NA							
		III	. CONCO	MITANT	DRUG	G(S)	AND H	IST	OF	۲Y											
22. CONCOMITANT DRU	UG(S) AND DATES OF ADM	1INISTRA	FION (exclude thos	se used to trea	t reaction)																
23. OTHER RELEVANT	HISTORY. (e.g. diagnostics,	allergies,	pregnancy with las	st month of pe	riod, etc.)																
From/To Dates Unknown to Ongo		Ту	pe of History / Not urrent Condi	es	Descrip		Obesity)														
	3	D	uration not re	eported		- , (,														
			D / B / C C C C C		1055		00111														
24a. NAME AND ADDRF	ESS OF MANUFACTURER		IV. MAN	UFACT		INF B. REMA		Ю	N								—	—			
Novo Nordisk A/S Lise Grimmeshave						Medically Confirmed: No															
Vandtaarnsvej 114 Soeborg, DK-2860, DENMARK																					
Phone: +45 44448																					
	04F MED 00	NITPO! *	10		05	5h NI^N*	IE AND ADD	000	05.5	EDO	DTF										
	24b. MFR CC		iO.				IE AND ADDF AND ADD														
24c. DATE RECEIVED																					
24c. DATE RECEIVED BY MANUFACTURE			LITERATUI	RE																	
30-JUN-2025	HEALTH		OTHER:																		
DATE OF THIS REPORT 23-JUL-2025		T TYPE		D.																	
	⊠ INITIAL		FOLLOWU	г.																	

Mfr. Control Number: 1472780

ADDITIONAL INFORMATION

7+13. DESCRIBE REACTION(S) continued

Patient's height: 160 cm.

Patient's weight: 120 kg.

Patient's BMI: 46.8750.

This non-serious Solicited Report from COSTA RICA was reported by a Consumer as "Headache(Headache)" with an unspecified onset date and concerned a 45 Years old Female patient who was treated with Saxenda (liraglutide 6 mg/mL) from unknown start date and ongoing for "Obesity",

Dosage Regimens:

Saxenda:

Current Condition: Obesity.

Batch Numbers: Saxenda: ASKU;

Action taken to Saxenda was reported as No Change.

The outcome for the event "Headache(Headache)" was Not recovered.

Reporter's causality (Saxenda) - Headache(Headache): Unknown

Company's causality (Saxenda) - Headache(Headache) : Possible