

## SUSPECT ADVERSE REACTION REPORT

## I. REACTION INFORMATION

1. PATIENT INITIALS (first, last) <b>PRIVACY</b>	1a. COUNTRY <b>COSTA RICA</b>	2. DATE OF BIRTH Day Month Year <b>PRIVACY</b>	2a. AGE <b>48</b> Years	3. SEX <b>Female</b>	3a. WEIGHT <b>99.00</b> kg	4-6 REACTION ONSET Day Month Year <b>JUL 2024</b>	8-12 CHECK ALL APPROPRIATE TO ADVERSE REACTION <input type="checkbox"/> PATIENT DIED <input checked="" type="checkbox"/> INVOLVED OR PROLONGED INPATIENT HOSPITALISATION <input type="checkbox"/> INVOLVED PERSISTENT OR SIGNIFICANT DISABILITY OR INCAPACITY <input type="checkbox"/> LIFE THREATENING <input type="checkbox"/> CONGENITAL ANOMALY <input checked="" type="checkbox"/> OTHER
7 + 13 DESCRIBE REACTION(S) (including relevant tests/lab data) Event Verbatim [PREFERRED TERM] (Related symptoms if any separated by commas) <b>Other Serious Criteria: Medically Significant</b> <b>chronic granulomatous mastitis [Plasma cell mastitis]</b> <b>rebound effect [Rebound effect]</b> <b>nausea [Nausea]</b> <b>stomach hurt a lot [Abdominal pain upper]</b> <b>sensitivity in the area of application [Injection site discomfort]</b> <b>disgust [Malaise]</b>  Case Description: This serious Solicited Report case from COSTA RICA was reported by a Consumer as "chronic granulomatous" <b>(Continued on Additional Information Page)</b>							

## II. SUSPECT DRUG(S) INFORMATION

14. SUSPECT DRUG(S) (include generic name) <b>#1 ) Saxenda (liraglutide 6 mg/mL) Solution for injection, 6 mg/mL</b> <b>(Continued on Additional Information Page)</b>	20. DID REACTION ABATE AFTER STOPPING DRUG? <input type="checkbox"/> YES <input type="checkbox"/> NO <input type="checkbox"/> NA
15. DAILY DOSE(S) <b>#1 ) 0.6 mg, qd</b>	16. ROUTE(S) OF ADMINISTRATION <b>#1 ) Subcutaneous</b>
17. INDICATION(S) FOR USE <b>#1 ) Obesity (Obesity)</b> <b>(Continued on Additional Information Page)</b>	21. DID REACTION REAPPEAR AFTER REINTRODUCTION? <input type="checkbox"/> YES <input type="checkbox"/> NO <input type="checkbox"/> NA
18. THERAPY DATES(from/to) <b>#1 ) JUL-2024 / DEC-2024</b>	19. THERAPY DURATION <b>#1 ) Unknown</b>

## III. CONCOMITANT DRUG(S) AND HISTORY

22. CONCOMITANT DRUG(S) AND DATES OF ADMINISTRATION (exclude those used to treat reaction) <b>#1 ) XIGDUO (DAPAGLIFLOZIN PROPANEDIOL MONOHYDRATE, METF</b> <b>#2 ) LEVOTHYROXINE (LEVOTHYROXINE) ; 2013 / Ongoing</b>  <b>(Continued on Additional Information Page)</b>
23. OTHER RELEVANT HISTORY. (e.g. diagnostics, allergies, pregnancy with last month of period, etc.) From/To Dates Type of History / Notes Description <b>Unknown to Ongoing Current Condition Obesity (Obesity)</b> <b>Unknown to Ongoing Current Condition Insulin resistance (Insulin resistance)</b>

## IV. MANUFACTURER INFORMATION

24a. NAME AND ADDRESS OF MANUFACTURER <b>Novo Nordisk A/S</b> <b>Lise Grimmeshave</b> <b>Vandtaarnsvej 114</b> <b>Soeborg, DK-2860 DENMARK</b> <b>Phone: +45 44448888</b>	26. REMARKS <b>Medically Confirmed: No</b>
24b. MFR CONTROL NO. <b>1469082</b>	25b. NAME AND ADDRESS OF REPORTER <b>NAME AND ADDRESS WITHHELD.</b>
24c. DATE RECEIVED BY MANUFACTURER <b>23-JUN-2025</b>	24d. REPORT SOURCE <input checked="" type="checkbox"/> STUDY <input type="checkbox"/> LITERATURE <input type="checkbox"/> HEALTH PROFESSIONAL <input type="checkbox"/> OTHER:
DATE OF THIS REPORT <b>02-JUL-2025</b>	25a. REPORT TYPE <input checked="" type="checkbox"/> INITIAL <input type="checkbox"/> FOLLOWUP:

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**ADDITIONAL INFORMATION****7+13. DESCRIBE REACTION(S) continued**

mastitis(Idiopathic granulomatous mastitis)" beginning on 07-APR-2025, "rebound effect(Rebound effect)" beginning on DEC-2024, "nausea(Nausea)" beginning on JUL-2024, "stomach hurt a lot(Stomach pain)" beginning on JUL-2024, "sensitivity in the area of application(Injection site discomfort)" beginning on JUL-2024, "disgust(Feeling unwell)" beginning on JUL-2024, and concerned a 48 Years old Female patient who was treated with Saxenda (liraglutide 6 mg/mL) from JUL-2024 to DEC-2024 for "Obesity", "insulin resistance",

Patient's height: 159 cm

Patient's weight: 99 kg

Patient's BMI: 39.15984340.

Dosage Regimens:

Saxenda: ??-JUL-2024 to ??-DEC-2024;

Current Condition: Obesity, Insulin resistance, Hypothyroidism, Diabetes mellitus.

Concomitant products included - XIGDUO(DAPAGLIFLOZIN PROPANEDIOL MONOHYDRATE, METFORMIN HYDROCHLORIDE), LEVOTHYROXINE

Since an unknown date in JUL-2024 had a lot of disgust, too many nausea, stomach hurt a lot, and she had sensitivity in the area after a certain time of application in the abdomen.

Since an unknown date in DEC-2024 patient stopped consuming the medication and the rebound effect was terrible, its a dreadful thing. She really reconsidered whether or not she should continue with a medication that, at the moment she stopped using it, the rebound effect was too much  
the rebound effect was brutal; from an unknown date in DEC or MAR (year unspecified), patient was at 103 kg, and was currently at 109 kg.

On 25-MAR-2025 patient felt her breast

Since 07-APR-2025 patient started having issues, developed an infection; on an unknown date underwent a biopsy (biopsy) (result unspecified)

On the same date a mammogram (mammogram) was done and , they told it was a tumor

Since 24-APR-2025 to 30-APR-2025 the patient was hospitalized due to the same

Patient already had the infection in her breast before, since her breast was so infected inside, they hospitalized.

On 24-APR-2025 , when they tried to perform the biopsy, but they could not do it because of the infection.

On 21-MAY-2025 They were able to perform the biopsy (result unspecified)

On 16-JUN-2025 the infected breast was diagnosed as benign, it was called chronic granulomatous mastitis

Batch Numbers:

Saxenda: has been requested

Action taken to Saxenda was reported as Product discontinued.

The outcome for the event "chronic granulomatous mastitis(Idiopathic granulomatous mastitis)" was Not Reported.

The outcome for the event "rebound effect(Rebound effect)" was Not recovered.

On DEC-2024 the outcome for the event "nausea(Nausea)" was Recovered.

On DEC-2024 the outcome for the event "stomach hurt a lot(Stomach pain)" was Recovered.

On DEC-2024 the outcome for the event "sensitivity in the area of application(Injection site discomfort)" was Recovered.

On DEC-2024 the outcome for the event "disgust(Feeling unwell)" was Recovered.

Company comment:

Idiopathic granulomatous mastitis is assessed as unlisted according to the Novo Nordisk company core data sheet (CCDS) for Saxenda

Information regarding circumstances leading to the reported event (such as autoimmune disorder, hormonal imbalance), social history (smoking) are unavailable which limits the medical assessment of the case. Recent infection of the breast can be considered as a risk factor for development of the reported event. Considering the pharmacological properties of the suspect product and nature of the event, idiopathic granulomatous mastitis is evaluated as unlikely related to the suspect product

**ADDITIONAL INFORMATION****7+13. DESCRIBE REACTION(S) continued**

This single case report is not considered to change the current knowledge of the safety profile of Saxenda

**13. Lab Data**

#	Date	Test / Assessment / Notes	Results	Normal High / Low
1		Biopsy		
		On an unknown date patient underwent a biopsy (result unspecified)		
2	21-MAY-2025	Biopsy		
		On 21-MAY-2025 patient underwent a biopsy (result unspecified)		
3	07-APR-2025	Mammogram		
		On 07-APR-2025 patient had a mammogram and they told it was a tumor		
4		Weight	109 kg	
5		Weight	103 kg	

**13. Relevant Tests**

from December to March (year unspecified), patient was at 103 kg,  
 On an unknown date patient underwent a biopsy (result unspecified)  
 On 21-MAY-2025 patient underwent a biopsy (result unspecified)  
 On 07-APR-2025 patient had a mammogram and they told it was a tumor

**14-19. SUSPECT DRUG(S) continued**

14. SUSPECT DRUG(S) (include generic name)	15. DAILY DOSE(S); 16. ROUTE(S) OF ADMIN	17. INDICATION(S) FOR USE	18. THERAPY DATES (from/to); 19. THERAPY DURATION
#1 ) Saxenda (liraglutide 6 mg/mL) Solution for injection, 6 mg/mL; Regimen #1	0.6 mg, qd; Subcutaneous	Obesity (Obesity) insulin resistance (Insulin resistance)	JUL-2024 / DEC-2024; Unknown

**22. CONCOMITANT DRUG(S) AND DATES OF ADMINISTRATION continued**

#1 ) XIGDUO (DAPAGLIFLOZIN PROPANEDIOL MONOHYDRATE, METFORMIN HYDROCHLORIDE) ; 2019 / Ongoing

**23. OTHER RELEVANT HISTORY continued**

From/To Dates	Type of History / Notes	Description
Unknown to Ongoing	Current Condition	Hypothyroidism (Hypothyroidism);
Unknown to Ongoing	Current Condition unknown type and duration	Diabetes (Diabetes mellitus);