

SUSPECT ADVERSE REACTION REPORT

I. REACTION INFORMATION

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| 1. PATIENT INITIALS (first, last) PRIVACY | 1a. COUNTRY COSTA RICA | 2. DATE OF BIRTH Day Month Year PRIVACY | | | 2a. AGE 34 Years | 3. SEX Female | 3a. WEIGHT 117.00 kg | 4-6 REACTION ONSET Day Month Year FEB 2024 | | | 8-12 CHECK ALL APPROPRIATE TO ADVERSE REACTION <input type="checkbox"/> PATIENT DIED <input 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) Other Serious Criteria: Medically Significant suicidal thoughts [Suicidal ideation] depression [Depression] panic attacks [Panic attack] anxiety increased [Anxiety] abrupt weight loss [Weight decreased] ozempic prescribed for weight loss [Off label use] patient indicates a dosage of 0.75 mg [Product use issue] Case Description: Study ID: 199-NovoDia (Continued on Additional Information Page) | | | | | | | | | | | |

II. SUSPECT DRUG(S) INFORMATION

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| 14. SUSPECT DRUG(S) (include generic name) #1) Semaglutide B 1.34 mg/ml PDS290 (SEMAGLUTIDE 1.34 mg/mL) Solution for injection, 1.34 mg/mL {Lot # (Continued on Additional Information Page) | | 20. DID REACTION ABATE AFTER STOPPING DRUG? <input type="checkbox"/> YES <input type="checkbox"/> NO <input checked="" type="checkbox"/> NA |
| 15. DAILY DOSE(S) #1) 0.25 mg, qw | 16. ROUTE(S) OF ADMINISTRATION #1) Subcutaneous | |
| 17. INDICATION(S) FOR USE #1) weight loss (Weight control) | | 21. DID REACTION REAPPEAR AFTER REINTRODUCTION? <input type="checkbox"/> YES <input type="checkbox"/> NO <input type="checkbox"/> NA |
| 18. THERAPY DATES(from/to) #1) FEB-2024 / Unknown | 19. THERAPY DURATION #1) Unknown | |

III. CONCOMITANT DRUG(S) AND HISTORY

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| 22. CONCOMITANT DRUG(S) AND DATES OF ADMINISTRATION (exclude those used to treat reaction) #1) ENALAPRIL (ENALAPRIL) Tablet ; Ongoing | | |
| 23. OTHER RELEVANT HISTORY. (e.g. diagnostics, allergies, pregnancy with last month of period, etc.) From/To Dates Type of History / Notes Description Unknown to Ongoing Current Condition Anxiety (Anxiety) Unknown to Ongoing Current Condition Blood pressure high (Hypertension) | | |

IV. MANUFACTURER INFORMATION

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| 24a. NAME AND ADDRESS OF MANUFACTURER Novo Nordisk A/S Lise Grimmeshave Vandtaarnsvej 114 Soeborg, DK-2860 DENMARK Phone: +45 44448888 | | 26. REMARKS Medically Confirmed: No World Wide #: CR-NOVOPROD-1417202 |
| | 24b. MFR CONTROL NO. 1417202 | 25b. NAME AND ADDRESS OF REPORTER NAME AND ADDRESS WITHHELD. |
| 24c. DATE RECEIVED BY MANUFACTURER 20-JUN-2025 | 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 26-JUN-2025 | 25a. REPORT TYPE <input type="checkbox"/> INITIAL <input checked="" type="checkbox"/> FOLLOWUP: 2 | |

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

Study description: Trial Title: Patient support programme to support physician and their daily work to maintain an optimal diabetic control of patients through added value services such as treatment starter kit, nutrition support through NovoDia call center, individual workshops, group workshops and free A1c test.

Patient's height: 159 cm.

Patient's weight: 117 kg.

Patient's BMI: 46.27981490.

This serious Solicited Report from COSTA RICA was reported by a Consumer as "suicidal thoughts(Suicidal ideation)" beginning on 2024 , "depression(Depression)" beginning on 2024 , "panic attacks(Panic attack)" beginning on 2024 , "anxiety increased(Anxiety aggravated)" beginning on 2024 , "abrupt weight loss(Weight loss)" beginning on 2024 , "ozempic prescribed for weight loss(Off label use)" beginning on FEB-2024 , "patient indicates a dosage of 0.75 mg(Unapproved dose administered)" with an unspecified onset date and concerned a 34 Years old Female patient who was treated with Ozempic (SEMAGLUTIDE 1.34 mg/mL) from FEB-2024 to MAY-2024 for "weight loss",

Dosage Regimens:

Ozempic: ??-FEB-2024 to Not Reported, ??-???-2024 to Not Reported, Not Reported to ??-MAY-2024;

Current Condition: Anxiety, High blood pressure

Historical Drug: Phentermine.

Concomitant medications included - ENALAPRIL.

Treatment medications included - FLUOXETINE, VENLAFAXINE.

On an unknown date in FEB-2024 the ozempic was prescribed for weight loss.

On an unknown date in 2024 the patient experienced depression to the point that she had to seek psychological help and take medications because she was having too many panic attacks. It was after about a month of using the medication that she started experiencing those symptoms, and it got worse when the dose was increased, so she decided to stop because she was not men-tally well anymore.

The patient also mentions that although she had anxiety issues before, the use of Ozempic was a trigger (referring to the significant increase in anxiety). She stated that she already knew there were those risks, but she didn't think she would experience them. She continued taking it be-cause she thought it might be temporary, but she reached the dose of 0.5 mg, and that's when she realized she was in a bad place because she literally had suicidal thoughts.

The patient reported that the Ozempic did work for her as she lost a lot of weight(Weight) but experienced a rebound effect afterward because of the depression. She started taking the medications, and they made her happy, and She started eating again' (referring to the rebound effect). The patient was still in a recovery state. It was also a very abrupt weight loss, her food intake de-created significantly.

It was reported that everything came back quadrupled, she don't know why, but she feels that injection really worsened her condition. She always told her psychologist 'The time she was thin-nest was the time she was saddest in her entire life,' and she states that despite the effects, she continued the medication until she finished it.

On an unknown date, the patient stated that patient started everything using Ozempic the panic attacks appeared with the use of Ozempic. The product works, but at least I imagine I was predisposed to anxiety, and that predisposition also contributed to the medication harming me. Instead of being happy about the weight loss, it was the darkest, saddest moment I felt. It works, but the person must be very mentally strong to handle that medication because it is too strong and it destroys your mind. If you are going to use it for weight loss, it destroys your mind. Patient mentions that she feels mentally weak. Additionally, she states that she has not had any changes in her diet and is trying to stay calmer

On an unknown date,the patient had taken a dosage of 0.75 mg.

On an unknown date,the patient mentioned that when discontinuing the medication, symptoms such as panic attacks and sadness gradually decreased until they disappeared.

Batch Numbers:

Ozempic: PP5K434, PP5K434, PP5K434;

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

Action taken to Ozempic was reported as Product discontinued.

The outcome for the event "suicidal thoughts(Suicidal ideation)" was Recovering/resolving.

The outcome for the event "depression(Depression)" was Recovering/resolving.

The outcome for the event "panic attacks(Panic attack)" was Recovered.

The outcome for the event "anxiety increased(Anxiety aggravated)" was Recovering/resolving.

The outcome for the event "abrupt weight loss(Weight loss)" was Recovering/resolving.

On MAY-2024 the outcome for the event "ozempic prescribed for weight loss(Off label use)" was Recovered.

On MAY-2024 the outcome for the event "patient indicates a dosage of 0.75 mg(Unapproved dose administered)" was Recovered.

Reporter's causality (Ozempic) -

suicidal thoughts(Suicidal ideation) : Possible

depression(Depression) : Possible

panic attacks(Panic attack) : Possible

anxiety increased(Anxiety aggravated) : Possible

abrupt weight loss(Weight loss) : Possible

ozempic prescribed for weight loss(Off label use) : Unknown

patient indicates a dosage of 0.75 mg(Unapproved dose administered) : Unknown

Company's causality (Ozempic) -

suicidal thoughts(Suicidal ideation) : Unlikely

depression(Depression) : Unlikely

panic attacks(Panic attack) : Unlikely

anxiety increased(Anxiety aggravated) : Unlikely

abrupt weight loss(Weight loss) : Possible

ozempic prescribed for weight loss(Off label use) : Possible

patient indicates a dosage of 0.75 mg(Unapproved dose administered) : Possible

Since last submission,the following information has been updated

0.75 mg dose was added in suspect

Outcome of panic attack was updated

Event "Unapproved dose administered" was added

Narrative updated accordingly

References included:

Reference Type: E2B Company Number

Reference ID#: CR-NOVOPROD-1417202

Reference Notes:

Company comment:

Suicidal ideation, depression, panic attack and anxiety are assessed as unlisted; weight decreased, off label use, unapproved dose administered are assessed as listed events according to the Novo Nordisk current CCDS information on Ozempic. The information on relevant medical history (on traumatic or stressful events, substance abuse) are not available. However, a medical history of anxiety is a confounder in this case. Hence the causality is assessed as unlikely for the events Suicidal ideation, depression, panic attack and anxiety; possible for events weight decreased, off label use, unapproved dose administered. This single case report is not considered to change the current knowledge of the safety profile of Ozempic.

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) Semaglutide B 1.34 mg/ml PDS290 (SEMAGLUTIDE 1.34 mg/mL) Solution for injection, 1.34 mg/mL {Lot # PP5K434; Exp.Dt. OCT-2026}; Regimen #1 | 0.25 mg, qw; Subcutaneous | weight loss (Weight control) | FEB-2024 / Unknown; Unknown |
| #1) Semaglutide B 1.34 mg/ml PDS290 (SEMAGLUTIDE 1.34 mg/mL) Solution for injection, 1.34 mg/mL {Lot # PP5K434; Exp.Dt. OCT-2026}; Regimen #2 | 0.5 mg, qw; Subcutaneous | weight loss (Weight control) | 2024 / Unknown; Unknown |
| #1) Semaglutide B 1.34 mg/ml PDS290 | 0.75 mg; Subcutaneous | weight loss (Weight control) | Unknown / MAY-2024; |

ADDITIONAL INFORMATION

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 |
|-----------------------------------------------------------------------------------------------------------|---------------------------------------------|---------------------------|------------------------------------------------------|
| (SEMAGLUTIDE 1.34 mg/mL) Solution for injection, 1.34 mg/mL {Lot # PP5K434; Exp.Dt. OCT-2026}; Regimen #3 | | | Unknown |

23. OTHER RELEVANT HISTORY continued

| From/To Dates | Type of History / Notes | Description |
|---------------|-------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Unknown | Historical Drug | Phentermine (PHENTERMINE); Drug Indication: Product used for unknown indication (Product used for unknown indication), Drug Reaction: No adverse reaction (No adverse event) |