

SUSPECT ADVERSE REACTION REPORT

I. REACTION INFORMATION

1. PATIENT INITIALS (first, last) PRIVACY	1a. COUNTRY COSTA RICA	2. DATE OF BIRTH			2a. AGE 67 Years	3. SEX Female	3a. WEIGHT Unk	4-6 REACTION ONSET			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
		Day	Month	Year				Day	Month	Year	
			PRIVACY					11	NOV	2024	

7 + 13 DESCRIBE REACTION(S) (including relevant tests/lab data)
 Event Verbatim [PREFERRED TERM] (Related symptoms if any separated by commas)
 fluid in right lung [Pulmonary oedema]
 Kidney test values were very altered [Renal disorder]
 Stomach pain [Abdominal pain upper]
 Swollen hands and legs from the knee to the fingers [Peripheral swelling]
 Swelling in her stomach/ stomach was inflamed/significant inflammation [Gastritis]
 Diarrhea [Diarrhoea]
 No energy [Asthenia]
 Not hungry [Decreased appetite]
 Diarrhea (second episode) [Diarrhoea]

(Continued on Additional Information Page)

II. SUSPECT DRUG(S) INFORMATION

14. SUSPECT DRUG(S) (include generic name) #1) Abemaciclib (Abemaciclib) Tablet		20. DID REACTION ABATE AFTER STOPPING DRUG? <input type="checkbox"/> YES <input checked="" type="checkbox"/> NO <input type="checkbox"/> NA
15. DAILY DOSE(S) #1) 150 mg, bid	16. ROUTE(S) OF ADMINISTRATION #1) Oral	
17. INDICATION(S) FOR USE #1) Breast cancer (Breast cancer)		21. DID REACTION REAPPEAR AFTER REINTRODUCTION? <input type="checkbox"/> YES <input type="checkbox"/> NO <input type="checkbox"/> NA
18. THERAPY DATES(from/to) #1) 08-NOV-2024 / 22-DEC-2024	19. THERAPY DURATION #1) 1 month 15 days	

III. CONCOMITANT DRUG(S) AND HISTORY

22. CONCOMITANT DRUG(S) AND DATES OF ADMINISTRATION (exclude those used to treat reaction) #1) ANASTROZOLE (ANASTROZOLE) Tablet ; MAY-2024 / Unknown #2) IRBESARTAN (IRBESARTAN) Unknown ; 2022 / Ongoing													
23. OTHER RELEVANT HISTORY. (e.g. diagnostics, allergies, pregnancy with last month of period, etc.) <table border="0"> <tr> <td>From/To Dates</td> <td>Type of History / Notes</td> <td>Description</td> </tr> <tr> <td>2022 to Ongoing</td> <td>Medical Condition</td> <td>Cataracts (Cataract)</td> </tr> <tr> <td></td> <td>her eyesight is failing. She indicates that with the chemotherapy her eyesight is getting worse</td> <td></td> </tr> <tr> <td>Unknown</td> <td>Medical Condition</td> <td>Oral intake reduced (Hypophagia)</td> </tr> </table>		From/To Dates	Type of History / Notes	Description	2022 to Ongoing	Medical Condition	Cataracts (Cataract)		her eyesight is failing. She indicates that with the chemotherapy her eyesight is getting worse		Unknown	Medical Condition	Oral intake reduced (Hypophagia)
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IV. MANUFACTURER INFORMATION

24a. NAME AND ADDRESS OF MANUFACTURER Eli Lilly Interamerica Inc (AR Branch) Tronador 4890 - Piso 12 Buenos Aires, Capital Federal CP: 1430 ARGENTINA Phone: 54 1145464000		26. REMARKS
	24b. MFR CONTROL NO. CR202411011684	
24c. DATE RECEIVED BY MANUFACTURER 24-MAR-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 27-MAR-2025	25a. REPORT TYPE <input checked="" type="checkbox"/> INITIAL <input type="checkbox"/> FOLLOWUP:	

25b. NAME AND ADDRESS OF REPORTER
NAME AND ADDRESS WITHHELD.

NAME AND ADDRESS WITHHELD.

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

Case Description: This solicited case reported by a consumer via a patient support program (PSP), with additional information from the initial reporting consumer via PSP from business partner, concerned a 67-year-old (at the time of initial report) female patient of an unknown origin.

Medical history included cataracts (she indicates that her eyesight was failing and that she had cataracts. She indicates that that had been going on for 2 years. She indicates that with the chemotherapy her eyesight was getting worse and that on 29-Nov-2024 she has an appointment with a doctor to see if she can have surgery), chemotherapy (she indicates that she started on 08-Dec-2023 and finished on 29-May-2024 and that since the chemotherapy she eats less), Diabetes, High blood pressure, thyroid problems (she did not know her diagnosis) and kidney test values were a little bad before starting abemaciclib therapy. Concomitant medications included milky insulin for diabetes, she was put on only that one because before she also had crystalline insulin, Thyroid pill (patient does not remember name), Irbesartan for blood pressure.

The patient received abemaciclib (Verzenio) tablets, 150 mg, twice daily, via orally for treatment of breast cancer beginning on 08-Nov-2024 or 09-Nov-2024 (conflicting date provided); also, she received anastrozole, 1 tablet daily as a concomitant medication for cancer since May-2024. On 11-Nov-2024, she had severe diarrhea and took two loperamide tablets and it stopped. Again on 14-Nov-2024 she had one diarrhea and took one loperamide pill and it stopped. On 29-Nov-2024 she was informed that her kidney values were very altered (no specific test, values, units or baseline were provided). Since an unknown date, she had experienced not having energy, was not hungry, stomach pain and diarrhea. On an unknown date in Dec-2024, she experienced ill and sick to her stomach, as her stomach was inflamed, she had swelling in both hands and legs from the knee to the fingers and she does not know the reason. As corrective treatment for stomach pain and reduce the swelling in her stomach she took an unspecified pill and for diarrhea she took loperamide, both symptoms improved temporarily. Due to her condition, she has not been using abemaciclib since 22-Dec-2024, because it caused significant inflammation and severe diarrhea. She used abemaciclib for approximately 22 days. She was currently at home. Next appointment with the oncologist would be in 6 months. Reportedly, she was hospitalized on 16-Mar-2025, because she had fluid in her right lung. She was not using abemaciclib therapy as she did not have it on site. Also, several tests were performed to her (unspecified). Further information regarding hospitalization and corrective treatments of the remaining events was not provided. Outcome of diarrhea (first episode) was recovered, for the events of lack of energy, decreased appetite, gastritis, diarrhea (second episode), pulmonary edema, and renal disorder was unknown and for the events of stomach pain and peripheral swelling was not recovered. The therapy status of abemaciclib therapy was drug discontinued.

The initial reporting consumer related the events of diarrhea and stomach inflammation with abemaciclib therapy and did not provide relatedness assessment of the rest of the events with abemaciclib therapy.

Update 26-Dec-2024: Additional information received on 18-Dec-2024 from the initial reporting consumer via PSP from business partner. Added events of lack of energy, decreased appetite, stomach pain, renal disorder and diarrhea (second episode). Added unspecified kidney test and renal disorder as medical history. Updated narrative with new information.

Update 28-Jan-2025: Additional information received on 21-Jan-2025 from the initial reporting consumer via PSP from business partner. Added two non-serious events peripheral swelling and gastritis. Updated suspect drug abemaciclib action taken from dose not changed to drug discontinued and outcome of the event stomach pain from unknown to not resolved and narrative with new information.

Update 19-Feb-2025: Additional information received on 13-Feb-2025 from the initial reporting consumer via PSP from business partner. Added the stop date of abemaciclib and added the severity of the event of diarrhea. Updated the causality for the events (diarrhea and stomach inflammation from not reported to yes). Updated narrative with new information.

Update 27-Mar-2025: Additional information was received from initial reporting consumer via PSP on 24-Mar-2025. This case was upgraded to serious upon addition of one serious event of pulmonary edema. Added one new laboratory test (unspecified). Updated narrative with new information.

13. Lab Data

#	Date	Test / Assessment / Notes	Results	Normal High / Low
1		Investigation		
		result not provided		
2		Renal function test		
		Reported as "very altered". No values, units or baseline were provided.		

ADDITIONAL INFORMATION

23. OTHER RELEVANT HISTORY continued

From/To Dates	Type of History / Notes	Description
Unknown	Medical Condition	Diabetes (Diabetes mellitus);
Unknown	Medical Condition	Blood pressure high (Hypertension);
Unknown	Medical Condition	Thyroid disorder (Thyroid disorder); she does not know her diagnosis
Unknown	Medical Condition	Renal disorder (Renal disorder);
08-DEC-2023 to 29-MAY-2024	Procedure	Chemotherapy (Chemotherapy); since the chemotherapy she eats less