

SUSPECT ADVERSE REACTION REPORT

I. REACTION INFORMATION

1. PATIENT INITIALS (first, last) PRIVACY	1a. COUNTRY COSTA RICA	2. DATE OF BIRTH			2a. AGE	3. SEX	3a. WEIGHT	4-6 REACTION ONSET			8-12 CHECK ALL APPROPRIATE TO ADVERSE REACTION
		Day	Month	Year	Unk	Male	Unk	Day	Month	Year	
		PRIVACY							Unk		
7 + 13 DESCRIBE REACTION(S) (including relevant tests/lab data)											
Event Verbatim [PREFERRED TERM] (Related symptoms if any separated by commas)		Product		Serious	Listed	Reporter Causality	Company Causality				<input type="checkbox"/> PATIENT DIED
Algo como alergia le molesta a paciente (not especifica) [Hypersensitivity]		XIGDUO		No	No	Related	Related				<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 type="checkbox"/> OTHER
(Continued on Additional Information Page)											

II. SUSPECT DRUG(S) INFORMATION

14. SUSPECT DRUG(S) (include generic name) #1) XIGDUO (DAPAGLIFLOZIN, METFORMIN) Tablet {Lot # Unknown}		20. DID REACTION ABATE AFTER STOPPING DRUG?
15. DAILY DOSE(S) #1) 5 milligram, bid	16. ROUTE(S) OF ADMINISTRATION #1) Oral use	<input type="checkbox"/> YES <input type="checkbox"/> NO <input type="checkbox"/> NA
17. INDICATION(S) FOR USE #1) Diabetes (Diabetes mellitus)		21. DID REACTION REAPPEAR AFTER REINTRODUCTION?
18. THERAPY DATES(from/to) #1) Unknown	19. THERAPY DURATION #1) Unknown	<input type="checkbox"/> YES <input type="checkbox"/> NO <input checked="" type="checkbox"/> NA

III. CONCOMITANT DRUG(S) AND HISTORY

22. CONCOMITANT DRUG(S) AND DATES OF ADMINISTRATION (exclude those used to treat reaction)		
23. OTHER RELEVANT HISTORY. (e.g. diagnostics, allergies, pregnancy with last month of period, etc.)		
From/To Dates Unknown to Ongoing	Type of History / Notes Indication	Description Diabetes (Diabetes)

IV. MANUFACTURER INFORMATION

24a. NAME AND ADDRESS OF MANUFACTURER AstraZeneca Serban Ghiorgiu 1 Medimmune Way Gaithersburg, Maryland 20878 UNITED STATES Phone: +1 301-398-0000		26. REMARKS World Wide #: CR-ASTRAZENECA-202507CAM009108CR Study ID: PSP-23269 Case References: CR-AstraZeneca-CH-00909474A
	24b. MFR CONTROL NO. 202507CAM009108CR	25b. NAME AND ADDRESS OF REPORTER NAME AND ADDRESS WITHHELD.
24c. DATE RECEIVED BY MANUFACTURER 11-JUL-2025	24d. REPORT SOURCE <input checked="" type="checkbox"/> STUDY <input type="checkbox"/> LITERATURE <input type="checkbox"/> HEALTH PROFESSIONAL <input type="checkbox"/> OTHER:	NAME AND ADDRESS WITHHELD.
DATE OF THIS REPORT 15-JUL-2025	25a. REPORT TYPE <input checked="" type="checkbox"/> INITIAL <input type="checkbox"/> FOLLOWUP:	

15-Jul-2025 21:10

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

Case Description: A solicited report has been received from a consumer in Patient Support Program. The report concerns a male patient born in 1947.

No medical history was reported.

No concomitant products were reported.

The patient started treatment with Xigduo (dapagliflozin, metformin) (batch number(s) Unknown) 5 milligram bid, Oral use, on an unknown date for diabetes.

On an unknown date, the patient experienced algo como alergia le molesta a paciente (not especifica) (preferred term: Hypersensitivity).

The outcome of the event(s) of algo como alergia le molesta a paciente (not especifica) was unknown.

The event was considered non-serious.

The reporter considered that there was a reasonable possibility of a causal relationship between Xigduo and the following event(s): algo como alergia le molesta a paciente (not especifica).

The company physician considered that there was a reasonable possibility of a causal relationship between Xigduo and the following event(s): algo como alergia le molesta a paciente (not especifica).