

SUSPECT ADVERSE REACTION REPORT	
BH-2025-016856	

I. REACTION INFORMATION

1. PATIENT INITIALS (first, last)	1a. COUNTRY	2. DATE OF BIRTH			2a. AGE Years	3. SEX	4-6 REACTION ONSET			8-12 CHECK ALL APPROPRIATE TO ADVERSE REACTION
PRIVACY	DOMINICAN	Day	Month	Year	Unkn	Male	Day	Month	Year	
	Cont..	??/??/??	??/??/??	??/??/??			??/??/??	??/??/??	??/??/??	
7+13 DESCRIBE REACTION(S) (including relevant tests/lab data)										

MedDRA Version : v.28.0
 1) DIARRHEA (Diarrhea (10012727), Diarrhoea (10012735))
 Recovered/Resolved

☐ PATIENT DIED
☐ LIFE THREATENING
☐ INVOLVED OR PROLONGED INPATIENT HOSPITALIZATION
☐ RESULTS IN PERSISTENCE OR SIGNIFICANT DISABILITY/INCAPACITY
☐ CONGENITAL ANOMALY
☐ OTHER MEDICALLY IMPORTANT CONDITION

II. SUSPECT DRUG(S) INFORMATION

14. SUSPECT DRUG(S)(include generic name)		20. DID EVENT ABATE AFTER STOPPING DRUG? <input type="checkbox"/> YES <input type="checkbox"/> NO <input checked="" type="checkbox"/> NA
1) CARDISPAN (LEVOCARNITINE) (Suspect) (Solution for injection)(Unknown)		
15. DAILY DOSE(S)		
Unkn		
16. ROUTE(S) OF ADMINISTRATION		21. DID EVENT REAPPEAR AFTER REINTRODUCTION <input type="checkbox"/> YES <input type="checkbox"/> NO <input checked="" type="checkbox"/> NA (NA : Not Applicable)
1) Unknown		
17. INDICATION(S) FOR USE		
1) DIALYSIS [10061105 - Dialysis]		
18. THERAPY DATE(S) (from/to)	19. THERAPY DURATION	
??/??/??	Unkn	

III. CONCOMITANT DRUG(S) AND HISTORY

22. CONCOMITANT DRUG(S) AND DATES OF ADMINISTRATION (exclude those used to treat reaction)
No concomitants used/reported
23. OTHER RELEVANT HISTORY (e.g. diagnostics, allergies, pregnancy with last month of period, etc.)
MedDRA Version : v.28.0
1) CHRONIC KIDNEY DISEASE (10064848, Chronic kidney disease) (Continuing: Yes)

IV. MANUFACTURER INFORMATION

24a. NAME AND ADDRESS OF MANUFACTURER		
Name : Bausch Health 400 Somerset Corporate Boulevard Bridgewater, NJ, UNITED STATES OF AMERICA Submission_ICSR_BHC@bauschhealth.com		
24. REPORT NULLIFIED	24b. MFR CONTROL NO.	
<input type="checkbox"/> YES <input type="checkbox"/> NO	BH-2025-016856	
24c. DATE RECEIVED BY MANUFACTURER	24d. REPORT SOURCE	
27/Aug/2025	<input type="checkbox"/> STUDY <input type="checkbox"/> LITERATURE <input checked="" type="checkbox"/> HEALTH PROFESSIONAL	
DATE OF THIS REPORT	25a. REPORT TYPE	
03/Sep/2025	<input checked="" type="checkbox"/> INITIAL <input type="checkbox"/> FOLLOWUP	

= Continuation attached sheet(s)..

Continuation Sheet for CIOMS report

1a. COUNTRY

DOMINICAN REPUBLIC

7+13 DESCRIBE REACTION(S) (including relevant tests/lab data) (Continuation...)

Event Description :

This non-serious spontaneous case was received on 27-Aug-2025 from a Physician via a sales representative and concerned a patient of unknown age and male gender.

Concurrent conditions: CHRONIC KIDNEY DISEASE MEDICATION

Medical history: Not Reported

Concomitant medications: Unspecified Chronic Kidney Disease medication

Past medications: Not Reported

Company suspect products include: CARDISPAN, Solution for injection (Batch number: unknown; Expiry Date: not reported) Start date not provided, Stop date not provided, administered via Unknown route for DIALYSIS.

On 27-Aug-2025 the patient experienced Diarrhea after administration of Cardispan (Levocarnitine) Injectable Solution. The treatment was not de-challenged; hence it was not changed to another drug. The patient did not receive any corrective treatment for adverse event. Concomitant treatment reported was unknown for Chronic Kidney Disease. By the time of the report, patient was recovered. No more information is available. This case is considered closed.

Reported Term (Preferred Term): Onset Date-Cessation Date; Seriousness; Outcome:
DIARRHEA(Diarrhoea): unknown - unknown; non-serious; Recovered/Resolved.

Action taken with company suspect product in response to the event, Dechallenge and Rechallenge:
CARDISPAN/DIARRHEA: Dose not changed; Not applicable; Not Applicable

Final therapy status: The patient was using Cardispan at the time of reporting.

Reporter's causality assessment:
CARDISPAN/DIARRHEA: Possible

Internal reference number included local PV number: IQF/GROSS/00006/2025.

This case is cross-referenced to DO-BAUSCH-BH-2025-016855 due to Same patient.

Company Remarks (Sender's Comments) :

Version 0 (27-Aug-2025):

The event Diarrhoea is assessed as non-serious. Diarrhoea is assessed as possibly related to CARDISPAN.

14.SUSPECT DRUG(S) (Continuation...)

Product-Reaction Level

1) Drug	: CARDISPAN
Active Substance	: LEVOCARNITINE
Drug Characterization	: Suspect
Form of Admin	: Solution for injection
Lot Number	: Unknown
Route of Admin	: Unknown
Indications	: DIALYSIS [10061105 - Dialysis]
Action(s) Taken With Drug	: Unknown

Causality

1) DIARRHEA (Diarrhea - 10012727, Diarrhoea - 10012735)	
Causality as per reporter	: Possible
Causality as per Mfr	: Possible
DeChallenge	: Not applicable

Continuation Sheet for CIOMS report

ReChallenge : Not Applicable

Labeling :

1) DIARRHEA
CORE

Labeled