

Mandatory Medical Device Problem Reporting Form for Industry

CANADA VIGILANCE - MEDICAL DEVICE PROBLEM REPORTING PROGRAM If more space is required, please attach additional sheets Fields required to be completed for updates/final reports are indicated by an *

					Page 1_	of ²
A. REPORTER INFORM	IATION					
1. i. Reporter Type Manufacturer Importer		3. Reporter File No. * INF-US-2020-030081 (412569)				
In the case where the reporter is the importer: ii. Did the importer report the incident to the manufacturer? O Yes No No iii. Is the importer also submitting the report on behalf of the		4. Health Canada File No. (if applicable) * 5. Type of Report * OPreliminary OUpdate OFinal OPreliminary & Final				
2. Reporter Contact Information *		(YYYY-MM-DD)				
Melissa Sanz, Quality Management Associate Becton Dickinson Canada Inc. 2100 Derry Road, Suite 100, Mississauga, ON L5N 0B3 tel: (905)288 -6148 / E-mail: Melissa.Sanz@bd.com		If "update/final", date the previous report was submitted to Health Canada: (YYYY-MM-DD) 6. Date Submitted * 2020-10-28 (YYYY-MM-DD)				
7. Name and Address	CareFusion 303, INC. 10020 Pacific Mesa Blvd. San Diego, California United States 92121-2733		Becton Dickinson Canada Inc. 2100 Derry Road, Suite 100, Mississauga, ON L5N 0B3			
8. Health Canada assigned company identification number (if known):	105302		101291			
9. Establishment License Number (if applicable):	Not Applicable		204			
B. INCIDENT INFORMA	ATION					
1. Classification of Incident * i. 010-Day 030-Day ii. 0 Canadian Foreign iii. Investigational testing Specia Radiation emitting device (if applic	5. Details of Incident It was reported that allegedly fourth top right and bottom left segments were dim for three devices and fourth top right segments were dim for two devices. Reportedly, the display boards of five large volume pump modules will be replaced.					
2. Date of Incident						
3. Reporter's Awareness Date 2020-10-08						
4. Patient Consequences						
There was no reported patient involvem	ent.					



	C. MEDICAL DEVICE INFORMATION
	Trade/Brand Name * laris Pump Module
	Control/Lot/Serial No. 4119327
3.	Expiration Date (YYYY-MM-DD)
4.	i. Device Classification I I I I I I I I I I I I I I I I I I I
5.	Software Version 9.1.17.7
6.	Age of Device 6 years 4 months
7.	How long was the device in use? Unknown
8.	Was the device labelled as sterile? Yes No
9.	Availability of device for evaluation Destroyed Returned to Manufacturer/Importer Neither (with explanation) No device will be returned per customer.
	D. COMPLAINANT INFORMATION Complainant is a:
2 .	Consumer OHealth professional Other Name of Complainant allum MacNicoll, Biomed
	Name of Health Care Facility (if applicable) BBOTSFORD REGIONAL HOSPITAL
32 Al	Address 2900 MARSHALL RD BBOTSFORD, BC, V2S 0C2 anada
+	Telephone No. and/or E-mail Address 1(604)851-4700 x642281 allum.macnicoll@fraserhealth.ca
l d a i i	Privacy Notice Statement: For the purposes of the Canada Vigilance - Medical Device Problem Reporting Program, information related to the identity of the complainant and/or reporter will be protected as personal information under the <i>Privacy Act</i> , and under the <i>Access to Information Act</i> in the case of an access to information request. For details with regard to personal information collected under this program, visit the Personal Information Bank; Health Canada; Health Products and Food Branch; Branch Incident Reporting System; HC PPU 088 at: http://infosource.gc.ca/inst/1476/1476-fedemp00-

eng.asp

E. INVESTIGATION INFORMATION

1. Investigative Actions and Timeline

The customer complaint could not be confirmed because the device was not returned for failure investigation.

This section only applies for preliminary & final, and final reports
2. Root Cause of Problem
The root cause of this failure was not identified.
3. Corrective Actions taken as a result of the investigation
This failure is being addressed through BD's corrective and preventive action processes.

Suthan, Thatpara (HC/SC)

From: CANDQualityCanada < CANDQualityCanada@bd.com>

Sent: 2020-10-28 2:53 PM

To: CANDQualityCanada; mdpr / dimm (HC/SC)

Subject: MDPR Health Canada Reports /

411003,411006,411196,412569,413178,413177,413179,413947

Attachments: 413177 Combined MDPR 28Oct2020.pdf; 412569 Combined MDPR 28Oct2020.pdf;

411196 Combined MDPR_28Oct2020.pdf; 411006 Combined MDPR_28Oct2020.pdf; 411003 Combined MDPR_28Oct2020.pdf; 413179 Combined MDPR_28Oct2020.pdf; 413178 Combined MDPR_28Oct2020.pdf; 413947 Combined MDPR_28Oct2020.pdf

Mandatory Problem Reporting - Completed Medical Device Problem Report from Becton Dickinson Canada Inc.

Please find enclosed, completed Medical Devices Problem Report Form for the following complaints:

BD Internal PIR File	Mandatory Report Type	Completed Forms
411003	30 day	Combined MPR
411006	30 day	Combined MPR
411196	30 day	Combined MPR
412569	30 day	Combined MPR
413178	30 day	Combined MPR
413177	30 day	Combined MPR
413179	30 day	Combined MPR
413947	30 day	Combined MPR

Abbreviations: PIR = Product Incident Report (Complaint), Combined = Preliminary and Final

We trust that you will find the present package satisfactory; however should you have any questions or concerns, please do not hesitate to contact me.

In the event that an email message is sent, we respectfully request that all of the aforementioned individuals be "cc'd" on the communiqué.

Thank you,





Melissa Sanz

Quality Associate
Quality Assurance

Melissa.Sanz@bd.com

Complaints: TOR-ComplaintsCA@bd.com

2100 Derry Rd W #100 Mississauga, ON L5N 0B3 Please note my number has changed:

Direct: +1.905.288.6148

Toll free: +1.800.268.5430 ext. 6148

<u>Learn more about BD.</u> Visit crbard.com



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Corporate Headquarters Mailing Address: BD (Becton, Dickinson and Company) 1 Becton Drive Franklin Lakes, NJ 07417 U.S.A.