

## 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 \*

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A. REPORTER INFORM	IATION				
1. i. Reporter Type  Manufacturer		3. Reporter File No. INF-US-2019-000085			
In the case where the reporter is the importer:		4. Health Canada File No. (if applicable) *			
ii. Did the importer report the incident to the manufacturer?  Yes  No  No  iii. Is the importer also submitting the report on behalf of the manufacturer?		5. Type of Report * OPreliminary OUpdate OFinal OPreliminary & Final			
O Yes O No		If "preliminary" only,	, anticipated date for th	he final report:	
2. Reporter Contact Information * b90a2a0b9b77ee4141076497accd53f8			e the previous report v	was submitted t	YYY-MM-DD)
				<u> </u>	YYY-MM-DD)
7. Name and Address	Manufactu	urer	Im	porter	
7. Name and Address	CareFusion Switzerland 317 Sarl  A-One Business Centre  Zone D'Activites  Vers-La-Piece No 10  Rolle, Switzerland, 1180  Becton Dickinson Canada Inc., 2100 Derry Road, Suite 100, Mississauga, Ontario Canada L5N 0B3		·		
8. Health Canada assigned company identification number (if known):	123422		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 0 Foreign iii. Investigational testing Specia Radiation emitting device (if applic	5. Details of Inciden	t			
2. Date of Incident	(YYYY-MM-DD)				
3. Reporter's Awareness Date	(YYYY-MM-DD)				
4. Patient Consequences		d41d8cd98f00b204e9	9800998ecf8427e		
60b725f10c9c85c70d97880dfe8191	b3				



C. MEDICAL DEVICE INFORMATION	E. INVESTIGATION INFORMATION
1. Trade/Brand Name * Extension set	Investigative Actions and Timeline
2. Control/Lot/Serial No.	-
3. Expiration Date	-
4. i. Device Classification  O   O    O    O    O     ii. Device License No.	
iii. Device Identification No	7ec31f1ef0a5435db98553ae4d91433b
iv. Manufacturer's Medical Device Identifier (catalogue/model no.) MZ5307	
5. Software Version	-
6. Age of Device	-
7. How long was the device in use?	This section only applies for preliminary & final, and final reports  2. Root Cause of Problem
8. Was the device labelled as sterile?  • Yes • No	-
9. Availability of device for evaluation  O Destroyed O Returned to Manufacturer/Importer  Neither (with explanation)	- 
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D. COMPLAINANT INFORMATION	I
1. Complainant is a: O Consumer O Health professional O Other	
<b>2. Name of Complainant</b> d41d8cd98f00b204e9800998ecf8427e	
3. Name of Health Care Facility (if applicable) d41d8cd98f00b204e9800998ecf8427e	Corrective Actions taken as a result of the investigation
4. Address	-
d41d8cd98f00b204e9800998ecf8427e	
5. Telephone No. and/or E-mail Address	
d41d8cd98f00b204e9800998ecf8427e	83cf4d005b3fd2ec0326b78c7d5c4a51
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	