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 $\ensuremath{^\star}$

			Page <u>1</u> of <u>2</u>				
A. REPORTER INFORMATION							
1. i. Reporter Type		3. Reporter File No. *					
O Manufacturer • Importer		PR-1626761-1634780					
In the case where the reporter is the importer: ii. Did the importer report the incident to the manufacturer? Yes No iii. Is the importer also submitting the report on behalf of the manufacturer? Yes No		4. Health Canada File No. (if applicable) * NOT AVAILABLE 5. Type of Report * ○ Preliminary ● Update ○ Final ○ Preliminary & Final If "preliminary" only, anticipated date for the final report: (YYYY-MM-DD) If "update/final", date the previous report was submitted to Health Canada: 2019-06-05 (YYYY-MM-DD)					
				2. Reporter Contact Information *			
				Baxter Corporation 7125 Mississauga Road Mississauga Ontario L5N0C2 Phone - 1(800) 387 8399 Fax - 1(855) 767			
						4572/Email: Randeep Cheema,	
				randeep_cheema@baxter.com		2020-07-16	(YYYY-MM-DD)
					Manufact	urer	Importer
7. Name and Address	BAXTER HEALTHCARE S Zurich, CH, 8010			SA Po Box 8010	BAXTER CORPORATION, 7125 MISSISSAUGA ROAD, MISSISSAUGA, ON, L5N 0C2		
8. Health Canada assigned	129789				101173		
company identification number							
(if known):							
9. Establishment License Number (if applicable):	N/A		191				
B INCIDENT INCORMATION							
B. INCIDENT INFORMATION		5.54.94.41.41	•				
1. Classification of Incident *		5. Details of Incident					
ii • Canadian • Service		On 11 Mar 2019, a healthcare professional reported Baxter					
II. ● Canadian		that three (3) EXACTAMIX EVA TPN BAG 2000ML (product code 740, lo# 60131932) had leaked at the port. The bags					
		were used to pump parenteral nutrition and leak was					
2. Date of Incident			administration to patient. Date of occurrence				
2019-02-12	(YYYY-MM-DD)	was on the 12th of February 2019. There was no patient involvement or medical injury involved.					
3. Reporter's Awareness Date		_ involvement or n	nedical injury involved.				
2019-03-11	(YYYY-MM-DD)						
	(1111-1010)	-					
4. Patient Consequences							
There was no patient involvement.							



C. MEDICAL DEVICE INFORMATION	E. INVESTIGATION INFORMATION
1. Trade/Brand Name * EXACTA-MIX EVA CONTAINER	1. Investigative Actions and Timeline
2. Control/Lot/Serial No.	This is being submitted as an update to the final mandatory report submitted on 2019-06-05.
3. Expiration Date	Two (2) samples were received at Baxter for evaluation.
S. Expiration Date (YYYY-MM-DD)	Functional testing was performed and revealed a leak
4. i. Device Classification	between the spike port tubing and spike port connector. The
\bigcirc I \bullet II \bigcirc III \bigcirc IV	reported issue was confirmed.
ii. Device License No.	A ball on the control for the control ball of the
5557	A batch review was performed for the reported Lot#, which indicated that this batch was released in accordance with
iii. Device Identification No 740	manufacturing specifications. No related defects or
iv. Manufacturer's Medical Device Identifier	exceptions were noted during the manufacturing, testing,
(catalogue/model no.)	and inspection of this lot.
H938740	
5. Software Version	
Not Applicable	
6. Age of Device Not Available	
7. How long was the device in use? UNKNOWN	This section only applies for preliminary & final, and final reports
8. Was the device labelled as sterile?	2. Root Cause of Problem
Yes O No	The cause was not determined, however the most likely
9. Availability of device for evaluation	cause of the reported condition was due to inadequate or
O Destroyed • Returned to Manufacturer/Importer	lack of cyclohexanone being applied to the cap when it was
O Neither (with explanation)	inserted to the spike port tubing during the manufacturing process,
Two samples were returned for evaluation.	process.
D. COMPLAINANT INFORMATION	
1. Complainant is a:	
O Consumer O Health professional ● Other	
2. Name of Complainant Kim Cook	
3. Name of Health Care Facility (if applicable)	
University of Alberta Hospital	
4. Address	3. Corrective Actions taken as a result of the investigation
8440-112 Street Edmonton AB T6G 2B7 CANADA	Baxter will continue to monitor this issue and any identified
	trend will be addressed by Corrective and Preventative
	Actions (CAPA) as necessary.
5. Telephone No. and/or E-mail Address	
780-407-8360/ Kim.Cook2@albertahealthservices.ca	
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

Matson, Sarah (HC/SC)

From: Canada SHS PS Quality <Canada_ProductSurveillance_Quality@baxter.com>

Sent: 2020-07-16 4:31 PM **To:** mdpr / dimm (HC/SC)

Subject: Four Final MDPRs updates PR-1626761-1664136 & PR-1626761-1634758 &

PR-1626761-1634746 & PR-1626761-1634780

Attachments: PR-1626761-1664136_Update to Final MDPR.pdf; PR-1626761-1634758_Update to Final

MDPR.pdf; PR-1626761-1634746_Update to Final MDPR.pdf; PR-1626761-1634780

_Update to Final MDPR.pdf

Dear Sir/Madam,

Please find enclosed four (4) follow-up MDPRs to four their respective Final Medical Device Problem Report submissions.

Please confirm receipt of this document. Thank you.

Christine He
Sr Quality Specialist, Product Surveillance
Baxter International Inc
7125 Mississauga Road
Mississauga, ON
L5N 0C2, Canada
Toll free 1-800-387-8399 x 6845

T: 905-369-6845 F: 1-855-767-4572