

URGENT: FIELD SAFETY NOTICE

Jelco® Optiva® IV Catheter 5063-Al

13th April 2023

Dear Valued Customers:

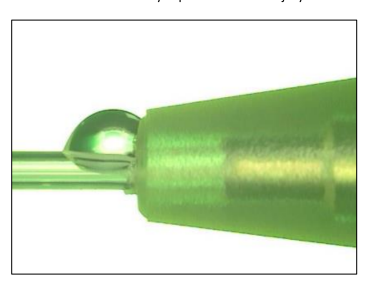
Smiths Medical, Inc. is issuing this Urgent Field Safety Notice to notify you of a potential defect with the Jelco® Optiva® V Catheters. This letter details the issue and the required steps for you to perform.

Issue:

Smiths Medical has identified the potential for a manufacturing defect within specific lots of the 24-gauge Jelco® Optiva® V catheters which may result in leakage at the insertion site.

Potential Risk:

The potential leakage is the result of catheter damage at proximal end of the catheter and near the tip of the female luer connector. A potential concern would be any fluid loss that would disrupt routine fluid delivery, drug delivery or administration of blood. To date, Smiths Medical has not received any reports of serious injury or death associated with this issue.



Affected Product:

Our records indicate that you may have received some of the affected products, which were distributed in United Kingdom in November and December 2022. The affected item and lot numbers are provided in Table 1, below:

Table 1: Affected Product and Lot Numbers

List Number	Product Description	Lot Number	
5063-AI	Jelco® Optiva® IV Catheter - 24G X 19mm	4331853 4321768	

Field Safety Notice: Optiva Catheters Potential Leakage

Smiths Medical Ref: FA2303-01



Required Actions for Users:

- 1) Please discontinue the use and distribution of the affected product immediately. Check your inventory and quarantine all affected product at your facility.
- 2) Inform potential users of the product in your organization of this notification and complete the attached response form, even if you do not have affected product. Please indicate on the form whether you intend to return this product to Smiths Medical or destroy it locally and return the completed response form to EMEA-Quality@icumed.com.
- 3) If you have distributed the product further, immediately notify your accounts that received the product identified in the Affected Product / Table 1 sections of this notification and ask your customers to complete a response form and return to you for overall completion.
- 4) Please contact Customer Service using the information provided below for assistance reordering replacement product.

Follow up Actions by Smiths Medical:

Upon receipt of the affected product, or upon receipt of a Certificate of Destruction, Smiths Medical will credit you for any product returned/destroyed. You will only receive credit for product that you return or that you certify has been destroyed locally. NOTE: Credits for product purchased through a distributor will be credited by the distributor.

For further inquiries, please contact Smiths Medical using the information provided below.

Smiths Medical Contact	Contact Information	Areas of Support	
Global Complaint Management	globalcomplaints@icumed.com	To report adverse events or product complaints	
Customer Support	ukcs@icumed.com	Additional information or assistance	

MHRA has been notified of this action.

Smiths Medical is committed to patient safety and is focused on providing exceptional product reliability and the highest level of customer satisfaction. Thank you for your prompt support on this important matter. We appreciate your cooperation.

Sincerely,

Joe Canavan Vice President, Quality, Consumables

Enclosures:

- Customer Response Form (pages 3 of this notice)
- Certificate of Destruction (separate file)

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URGENT: FIELD SAFETY NOTICE – RESPONSE FORM

Jelco® Optiva® IV Catheter 5063-AI

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Check your inventory and complete the information below, even if you do not have the affected product. <u>Failure to complete all sections of this page</u> may result in improper, delayed or denied credit.

Please return the completed form to EMEA-Quality@icumed.com. Smiths Medical Customer Service and your local sales representative.

Name of Hospital / Facility				
Hospital / Facility Address				
elephone Number				
ame and Title of Person Completing this Form				
ignature of Person Completing this Form				
ate				
Purchased through a distributor, please list distributor ame/location here for traceability purposes				
I have <u>NO</u> affected products (con YES, I have affected products If you have affected product TABLE 1			e)	
List Number	Lot Number	Quantity in inventory	PO, debit memo or invoice	
If you have distributed the respond to ICU Medical with TABLE 2			ated information received from you	ur custome rs
List Number	Lot Number	Quantity destroyed locally by customer	Quantity returned to distributor	
			ucts on site (complete and return p	rovided Certif
of Destruction to the email I have followed the inst return the affected product I have followed the inst	ructions provided to m s.	e and I will contact my Smiths N	Medical CS Representative to make	arrange ment s

Adverse events and complaints associated with the use of this product should be reported and emailed to the Competent Authority or to Smiths Medical's Global Complaint Management Department at globalcomplaints@icumed.com.

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