07.07.2011

URGENT FIELD SAFETY NOTICE

Commercial Name of Affected Product:	Ureter Catheter, curved, coaxial with no eyes		
Type of action:	Market Withdrawal		
Part Number:	264810-000040 20	64810-000050	264811-000040
I	134100-000040 20	64100-000050	264811-000050
Lot Number:		All lots	

Dear Customer,

1. Details of affected devices

Teleflex has issued an **extension** to the voluntary Field Action for products as detailed in above table. *All lots from the above part numbers are now affected.*

2. Description of the problem

Teleflex have received a small number of complaints that the white flexible sleeve/tip became detached from the metal stylet during use. The detached white flexible tip may remain in the patient resulting in potential injury and the need for possible medical intervention.

3. FIELD SAFETY CORRECTIVE ACTION INSTRUCTIONS:

Advice on action to be taken by Medical Staff

- 1. We request that you check stock for product within the scope of this field action. Users should cease use and distribution of stock from the affected lot number and quarantine all products from the affected lot number immediately.
- If you have no stock from the affected lot number, please indicate so on the 'field safety corrective action acknowledgement and stock status form' and fax to your local Teleflex Customer Service/Sales Rep as indicated on the 'field safety corrective action acknowledgement and stock status form' (Appendix 1)
- 3. If you do have stock from the affected lot number, please contact your local Teleflex Customer Service/Sales Rep (or Distributor) as indicated on the attached 'field safety corrective action acknowledgement and stock status form' for a Return Authorization Number. Once you have received the Return Authorization Number, please enter it in the space provided on the enclosed 'field safety corrective action acknowledgement and stock status form'
- 4. Complete the enclosed 'field safety corrective action acknowledgement and stock status form' providing detail on stock from the affected lot number under your control within the scope of this Field Safety Action and immediately email the completed form to <u>orders.intl@teleflexmedical.com</u>This will allow us to document your receipt of this letter and the amount of product you have on hand for return.
- 5. Your local Teleflex Customer Service/Sales Rep (or Distributor) will coordinate the product return with you.
- 6. Teleflex Customer Service can credit your account when the product is returned.

Instruction for Distributors of affected product

If you are a distributor, Teleflex requires that you communicate this field safety corrective action notice to your customers who received product within the scope of this field safety corrective action by providing:

- A copy of this Field Safety Notice to them.
- A copy of the 'field safety corrective action acknowledgement and stock status form' (Appendix 1)

The 'field safety corrective action acknowledgement and stock status form' is required to be completed in its entirety, signed and returned to you (the Distributor).

As a Distributor, it is your responsibility to provide Teleflex with confirmation that all of your consignees have been contacted under this field safety corrective action. Please forward the completed Acknowledgement Form to <u>orders.intl@teleflexmedical.com</u> or by fax on +353 1 437 0773

Please be advised that all affected European Economic Area/Switzerland (EEA/CH) Member State Competent Authorities will be notified by Teleflex where we have distributed product directly. If you are a distributor and/or have a reporting responsibility within or outside the EEA/CH area please notify your local Competent Authority of this action. Please forward the notification and all communication with your local competent authority to Teleflex.

4. Teleflex

Teleflex is notifying any potentially affected Customers, Teleflex employees and Distributors of this Field Safety Notice.

5. Transmission of this Field Safety Notice

This notice should be passed on all persons who need to be aware within your organization or to any organization where the potentially affected devices have been transferred.

6. Contact reference person

Should you require any further information or support concerning this issue, please contact:

For Customer Service Raychel Murtagh Email: <u>orders.intl@teleflexmedical.com</u> Fax: +353 1 437 0773 Phone: +353 906 460 838

For Product Specific Queries:

Vladamir Vasseck on Phone: +420 602 791 683 Email: vvasek@teleflexmedical.com

This field safety corrective action is voluntary and all affected competent authorities have been advised of this Field Safety Corrective Action.

Teleflex is committed to providing high quality, safe and effective products. We sincerely apologize for any inconvenience this action may cause your operations. If you have any other questions, feel free to contact your local sales representative or Customer Service at orders.intl@teleflexmedical.com

Signed by:



Attachments as follows: Appendix 1 :'field safety corrective action acknowledgement and stock status form'

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Appendix 1

FIELD SAFETY CORRECTIVE ACTION ACKNOWLEDGEMENT AND STOCK STATUS FORM

Medical Device Field Safety Corrective Action Concerning TeleflexProducts

Customer No	Immediate Attention Required RETURN COMPLETED FORM IMMEDIATELY TO:		Return Authorisation No	
Email: <u>orders.intl@teleflexmedical.com</u> ; Fax: +353 1 437 0773				
We have no inventory of stock from the affected lot number within the scope of this Field		We have the following affected stock from the affected lot number at our facility and have discontinued use and distribution. We have quarantined the affected product, and will return the following quantities.		
Action.		Please credit our account once you have received the returns		

Please print product numbers clearly.

Please credit our account once you have received the returns

Product	Ureter Catheter, curved, coaxial with no eyes (White sleeve)
Product Number	Lot Number/Quantity (unit)
134100-000040	
264100-000050	
264811-000040	
264810-000050	
264810-000040	
264811-000050	

Print Name/Title

Signature

Institution Name

Address

City, State, Zip Code

Date

Telephone Number