Smith & Nephew, Inc. 1450 Brooks Road Memphis, TN 38116 USA 1-901-396-2121 1-800-821-5700 www.smith-nephew.com



# Urgent Medical Device Recall Notice R-2018-06

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#### <Insert Address>

This letter is to inform you that Smith & Nephew Inc., have initiated a field action to voluntarily remove a single lot of PERI-LOC K-WIRE 1.6MM X 150MM LENGTH TROCAR POINT due to a labeling error. The incorrect expiration date of 21 July 2017 was provided on both the carton and pouch labels. The correct expiration date is 19 July 2027.

### Please see product details below:

Product Number	Description	Batch Number
71161016	PERI-LOC K-WIRE 1.6MM X 150MM LENGTH TROCAR POINT	17GK00029

Shipment Date: October 9, 2017 through December 11, 2017

## Potential Risk with Use of the Product

In the event the affected device is presented for use, the device will perform as indicated. Therefore use of or exposure to the affected device presents a negligible risk of adverse health consequences.

#### **Required Actions:**

• Please follow the instructions on the attached Response Form.

Enclosure: Response Form



# Urgent Medical Device Recall Notice R-2018-06

March 9, 2018 <Insert Address>

#### PLEASE COMPLETE ALL ITEMS AND RETURN WITHIN 5 DAYS OF RECEIPT

#### **Required Actions:**

- 1. Please inspect your inventory and locate any unused devices from the listed product and batch numbers on the first page of this Field Action Notification, and quarantine them immediately.
  - a. If you are a distributor, you must notify your customers of the field action and ensure that these actions are carried out.
- 2. If you have no product to return, please put an X in the appropriate location below.
- 3. If you have product to return, please list the batch numbers and quantities of each batch that you are returning in the appropriate boxes below.
- 4. Complete the remainders of the form sign and send to <u>FieldActions@smith-nephew.com</u> or fax to 901-566-7975.
  - **Please Note** even if you have no product to return, this form must be completed, signed and returned.
- 5. Once the form is received by Smith & Nephew, you will be sent a Return Authorization (RA) number.

If you have any questions or concerns regarding this recall please contact <a href="mailto:FieldActions@smith-nephew.com">FieldActions@smith-nephew.com</a>.

Product Part Number	Batch Number (List Specific Batch #'s to be Returned)	Quantity of Units to be Returned
•	of this Medical Device Field Action and it ha	s been communicated within
rganization.		
Printed Name (required):	Title:	
,	Title:[	
Signature (required):		Date (required):/_