URGENT: VOLUNTARY MEDICAL DEVICE RECALL
(REMOVAL)
5mm ENDOPATH® XCEL™ Trocars with Optiview Technology
(Product Codes 2B5LT, 2CB5LT, 2B5ST, 2CB5ST, 2B5XT)

[Date]

Dear Distributor:

Our records indicate that you have ordered 5mm ENDOPATH® XCEL™ Bladeless Trocars with Optiview Technology and may have received the product lots subject to this recall. PLEASE DISTRIBUTE THIS INFORMATION TO ALL PERSONNEL RESPONSIBLE FOR 5mm ENDOPATH® XCEL™ Trocars with Optiview Technology.

At Ethicon Endo-Surgery, LLC (“Ethicon”), our first priority is to support the needs of our customers and their patients, and that includes the safe and effective use of our products. Recently, Ethicon has become aware of reports from customers related to difficulty removing the obturator from the 5mm ENDOPATH® XCEL™ Trocars with Optiview Technology. **For that reason, we are issuing a voluntary recall of the following products with expiration dates between September 2022 through January of 2023.**

There have been no reported adverse events associated with this issue and it represents a low risk to patient health.

**EFFECTIVE IMMEDIATELY – DO NOT USE OR DISTRIBUTE THE FOLLOWING 5mm ENDOPATH® XCEL™ TROCARS WITH OPTIVIEW TECHNOLOGY. THIS RECALL DOES NOT AFFECT ANY OTHER PRODUCT CODES OR Lots WITH DIFFERENT EXPIRATION DATES.**

<table>
<thead>
<tr>
<th>PRODUCT NAME</th>
<th>PRODUCT CODE</th>
<th>Expiration Date Range</th>
<th>DESCRIPTION / SIZE</th>
</tr>
</thead>
<tbody>
<tr>
<td>5mm ENDOPATH® XCEL™ Bladeless Trocars with Optiview Technology</td>
<td>2B5ST</td>
<td>2022-10-31 through 2023-01-31</td>
<td>Stability sleeve, 5mm diameter (Length 75mm)</td>
</tr>
<tr>
<td></td>
<td>2B5LT</td>
<td>2022-09-30 through 2023-01-31</td>
<td>Stability sleeve, 5mm diameter (Length 100mm)</td>
</tr>
<tr>
<td></td>
<td>2B5XT</td>
<td>2022-11-30 through 2023-01-31</td>
<td>Stability sleeve, 5mm diameter (Length 150mm)</td>
</tr>
<tr>
<td>5mm ENDOPATH® XCEL™ Universal Sleeve Trocars with Optiview Technology</td>
<td>2CB5ST</td>
<td>2022-10-31 through 2023-01-31</td>
<td>Stability sleeve, 5mm diameter (Length 75mm)</td>
</tr>
<tr>
<td></td>
<td>2CB5LT</td>
<td>2022-09-30 through 2023-01-31</td>
<td>Stability sleeve, 5mm diameter (Length 100mm)</td>
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Table 2 has kit codes and lot numbers that contained trocars subject to this recall.

Table 2. Kit Codes with Trocars Lots Subject to this notification

<table>
<thead>
<tr>
<th>Kit Code</th>
<th>Kit Lot Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>LSR278</td>
<td>10064703</td>
</tr>
<tr>
<td>LSR278</td>
<td>10064715</td>
</tr>
<tr>
<td>LSR278</td>
<td>10064728</td>
</tr>
<tr>
<td>LSR278</td>
<td>10064749</td>
</tr>
</tbody>
</table>

Since there is a low risk to patient health due to this issue, health care practitioners who have treated patients using the 5mm ENDOPATH® XCEL™ Trocars with Optiview Technology should follow those patients post-operatively in the usual manner with no additional action required.

Refer to Attachment 1 for assistance in identifying the product lot subject to this recall.
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IDENTIFICATION OF THE PRODUCT LOT SUBJECT TO THIS RECALL:

The products subject to this recall in your inventory can be identified by product code and expiration date (See Table 1 above). The product code and expiration dates can be determined by using the Product Identification Tool within Attachment 1.

ACTION REQUIRED:

1. Examine your inventory immediately to determine if you have the product lot subject to this recall on hand and quarantine such product(s).

2. Remove the product lot subject to this recall and communicate the issue to all relevant materials management personnel, or anyone else in your facility who needs to be informed.

3. It is not necessary to contact your accounts regarding this recall. Ethicon will directly communicate with customers about the above listed product lot subject to this recall.

4. Complete the Business Reply Form (BRF) (Attachment 2) confirming receipt of this notice and fax or email it to [Insert Affiliate Information] within three (3) business days. Please return the BRF even if you do not have the product lot subject to this recall.

5. Customers are required to return all unused 5mm ENDOPATH® XCEL™ Trocars with Optiview Technology subject to this recall that are in their inventory immediately. Only unused 5mm ENDOPATH® XCEL™ Trocars with Optiview Technology subject to this recall returned by July 30, 2018 will be eligible for reimbursement. Any unused 5mm ENDOPATH® XCEL™ Trocars with Optiview Technology subject to this recall returned after July 30, 2018 will not be eligible for reimbursement.

6. To return unused 5mm ENDOPATH® XCEL™ Trocars with Optiview Technology subject to this recall, photocopy the completed BRF, place it in the box with the subject product(s), and affix the pre-paid authorized shipping label included with this recall notification letter. Ethicon will pay for the shipping charges only if the authorized label is used. Extra shipping labels may be obtained by calling [Insert Affiliate Information]. Your JJHCS number and mailing address have been pre-populated on the BRF.

If you require any assistance with returning product, subject to this recall, please contact [Insert Affiliate Information]. [Insert Affiliate Information] business hours are [Insert Affiliate Information].

We recognize disruptive to your facility and we apologize for any inconvenience this may cause.

If you have additional questions regarding this recall or to report any customer complaints, please contact [Insert Affiliate Information].
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As with any medical device, adverse reactions or quality problems experienced with the use of this product should be reported to your Sales Representative, directly to Ethicon, or your National Health Authority. If you have any further questions related to this notice or if you need any additional communications, please contact your local Sales Representative.

Attachments:
Attachment 1: Product Identification Tool
Attachment 2: Business Reply Form
URGENT: VOLUNTARY MEDICAL DEVICE RECALL (REMOVAL)
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ATTACHMENT 1: Product Identification Tool for 5mm ENDOPATH® XCEL™ Trocars with Optiview Technology (See Table 1 for affected product codes and lots.)

This tool will help customers identify the product lot of 5mm ENDOPATH® XCEL™ Trocars with Optiview Technology subject to this recall. Please refer to the table above for the product expiration dates subject to this recall.

SALES UNIT / DISPENSER CARTON (CONTAINING SIX (6) Trocars)

TOP OF SALES UNIT (WITH LOT AND EXPIRY DATE)
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TYVEK POUCH (CONTAINING ONE (1) TROCAR)

Lot Number
Expiration Date
Product Code
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ATTACHMENT 2: Business Reply Form (BRF)

Your timely response to this customer notification is requested. Please complete and fax this form to [Insert Affiliate Information] within 3 business days, even if you do not have the product lot subject to this recall to return.

If you have the devices from the product lot subject to this recall to return, please make a photocopy of your completed Business Reply Form and enclose with your return. Thank you for your cooperation.

Product Inventory – Please check one:

☐ We have NO remaining 5mm ENDOPATH® XCEL™ Trocars with Optiview Technology or kits subject to this recall.

☐ We have 5mm ENDOPATH® XCEL™ Trocars with Optiview Technology or kits subject to this recall and are returning the following products (Expiration Dates between September 2022 and January 2023):

<table>
<thead>
<tr>
<th>PRODUCT NAME</th>
<th>PRODUCT CODE</th>
<th>LOT #1</th>
<th>Quantity Returning (Eaches)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5mm ENDOPATH® XCEL™ Bladeless Trocars with Optiview Technology</td>
<td>2B5ST</td>
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Note¹: Please include lot number and not expiration date when completing this table.
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</tbody>
</table>

[Account Name]
[Account Address]

Print Name of Person Completing Business Reply Form:  
Telephone Number:

Account Number:  
(number used to order J&J product)  
Date:

Reimbursement Shipping Address (If different from above):

Signed*:  
*Your signature provides confirmation that you have received and understood this notification

Your comments are welcome.