

Cook Medical Europe

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Urgent Field Safety Notice

Commercial name of the affected product: EchoTip® Ultra Endobronchial High Definition Ultrasound

Needles

Manufacturer: Cook Ireland Ltd. FSCA-identifier: 2015FA0009
Type of action: IFU update

Date: 25 November 2015

Attention: Risk Management/Recall Administration

Details on affected devices:

Part Numbers:

ECHO-HD-22-EBUS-O (EchoTip® Ultra Endobronchial High Definition Ultrasound Needles) ECHO-HD-25-EBUS-O (EchoTip® Ultra Endobronchial High Definition Ultrasound Needles)

Affected Lot Numbers:

| Lot Number |
|------------|------------|------------|------------|------------|------------|
| C1116535 | C1126058 | C1143273 | C1152641 | C1163713 | C1170086 |
| C1116538 | C1127370 | C1143851 | C1152642 | C1164362 | C1170087 |
| C1116541 | C1130174 | C1144715 | C1152643 | C1164365 | C1172089 |
| C1117061 | C1130179 | C1146104 | C1152990 | C1164392 | C1173025 |
| C1117086 | C1130185 | C1146107 | C1154442 | C1164420 | C1174704 |
| C1117651 | C1131652 | C1146456 | C1155266 | C1164827 | C1174705 |
| C1119784 | C1135485 | C1148260 | C1156256 | C1167098 | |
| C1119788 | C1135487 | C1149128 | C1156856 | C1167549 | |
| C1119789 | C1138250 | C1149808 | C1156857 | C1167931 | |
| C1120397 | C1138252 | C1150868 | C1157014 | C1169098 | |
| C1120638 | C1139574 | C1152269 | C1158248 | C1169697 | |
| C1121308 | C1139576 | C1152271 | C1159099 | C1169698 | |
| C1121816 | C1139577 | C1152272 | C1159609 | C1169699 | |
| C1122234 | C1139585 | C1152273 | C1160628 | C1169919 | |
| C1124514 | C1139596 | C1152274 | C1161960 | C1169920 | |
| C1124521 | C1140105 | C1152468 | C1162430 | C1170082 | |
| C1125439 | C1140598 | C1152469 | C1163388 | C1170084 | |

Form: F14-00A (R7, CR15-0637)

Description of the problem:

Cook Ireland Ltd is updating the Instructions for Use (IFU) IFU0051 from revision 5 to revision 6 for the EchoTip® Ultra Endobronchial High Definition Ultrasound Needles to re-insert the following text "This device is designed for single use only. Attempts to reprocess, resterilize, and/or reuse may lead to device failure and/or transmission of disease" due to an omission of this text from Revision 5 of this IFU. This statement was included in the previous revision of the IFU, this notice is for customers who had not previously received the device and therefore may not be aware of the risks of re-use of this device as required by the MDD 93/42/EC.

This Field Safety Corrective Action is to correct the IFU; the IFU is being updated to reflect these changes. Neither device nor IFU need to be returned to Cook Medical.

Modifications made to the IFU are as follows:

The IFU is being updated to re-insert the following text:

"This device is designed for single use only. Attempts to reprocess, resterilize, and/or reuse may lead to device failure and/or transmission of disease"

Action to be taken by the user:

- 1. Please review the the attached list of affected products and lot numbers that were shipped to your account and identify any devices you still have in stock.
- 2. Please affix a copy of this Field Safety Notice to each of the affected products still in stock at your facility to ensure all users of these products are aware of the IFU updates.
- 3. Please note this it to inform you of the updates to the IFU of the affected products and there is no need to return any affected devices or IFUs.
- 4. Please complete the enclosed Customer Response Form and send via email to European.FieldAction@CookMedical.com or alternatively by fax to Cook Medical marked for the attention of European Customer Quality Assurance (fax number +353 61334441).

Transmission of this Field Safety Notice:

- This notice needs to be passed on to all those who need to be aware within your organisation or to any organisation or patient where the potentially affected devices have been transferred.
- Please transfer this notice to other organisations on which this action has an impact.
- Please maintain awareness on this notice and resulting action for an appropriate period to ensure effectiveness of the corrective action.

Contact reference person:

Sinead Burke
Director of Regulatory Affairs
COOK Ireland Ltd.
O'Halloran Road, National Technology Park, Limerick, IRELAND

Or

Annemarie Beglin Quality Systems Manager COOK Medical Europe O'Halloran Road, National Technology Park, Limerick, IRELAND

Should you have any questions, please feel free to contact us for more information (e-mail: European.FieldAction@CookMedical.com, phone +353 61 334440).

We confirm that this notice has been notified to the appropriate Regulatory Agency.

Signature

Annemarie Beglin

Annemarie Beglin Quality Systems Manager