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Rev 1: September 2018

FSN Ref: 385183-A

Date: 01/12/2022

Urgent Field Safety Notice

NEONATAL, NASAL CANNULA WITH CURVED PRONGS AND TUBE, 2.1M – REF: 1164000

PLEASE NOTE: THIS NOTICE SUPERCEDES THE PREVIOUS FIELD SAFETY NOTICE 385183 ISSUED ON 23/11/2022

For Attention of: All clinical staff, Managers and users of the above product

Contact details of local representative (name, e-mail, telephone, address etc.)*

Giedrius Budrys

Customer Resolution and Relationship Manager

Intersurgical UAB

Arnioniu str 60, LT-18170 Pabrade Lithuania

Email: giedriusb@intersurgical.lt

Tel. +370 387 66611

or

This could be a distributor or local branch of the manufacturer. To be added at the appropriate stage in the different local languages

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NEONATAL, NASAL CANNULA WITH CURVED PRONGS AND TUBE, 2.1M - REF: 1164000 Risk addressed by FSN

| | 1. Information on Affected Devices* |
|---|---|
| 1 | 1. Device Type(s)* |
| • | Basic Nasal Oxygen Cannula |
| 1 | 2. Commercial name(s) |
| | NEONATAL, NASAL CANNULA WITH CURVED PRONGS AND |
| | TUBE, 2.1M |
| 1 | 3. Unique Device Identifier(s) (UDI-DI) |
| | N/A |
| 1 | 4. Primary clinical purpose of device(s)* |
| | To deliver oxygen into a patient's nose. |
| 1 | 5. Device Model/Catalogue/part number(s)* |
| | 1164000 |
| 1 | 6. Software version |
| - | N/A |
| 1 | 7. Affected serial or lot number range |
| | 32101838 32103910 32104363 32105886 32107401 32107969 32108492 32108661 |
| | 32113543 32114272 32115031 32115516 32116417 32117131 32118980 32121015 |
| | 32121426 32122791 32200213 32201328 32203136 32204165 32205503 32205602 |
| | 32207055 32208628 32209416 32210009 32210465 32210778 32211422 32213464 |
| | 32214035 32214784 32215283 32215951 32217487 32218056 |
| 1 | 8. Associated devices |
| | N/A |
| | |

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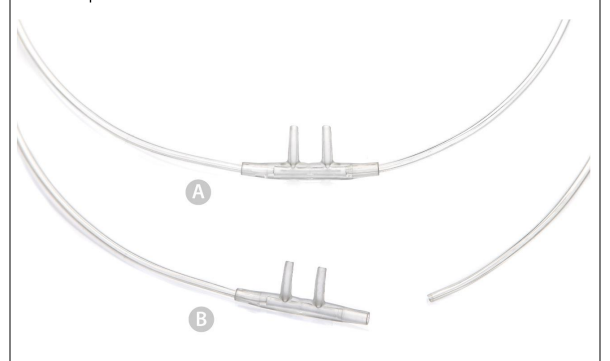
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2 Reason for Field Safety Corrective Action (FSCA)*

2. 1. Description of the product problem*

We have received reports related to disconnection of the tube from the nasal prong section while using our Neonatal Nasal Cannula with curved prongs and tube. Image A, below, shows the correct configuration, and image B demonstrates the reported disconnection.

The purpose of this FSN is to advise you that Intersurgical is issuing a Safety Notice for the removal of the potentially affected products. This safety notice applies to all distributed products with the Lot Numbers indicated above



2 2. Hazard giving rise to the FSCA*

If the oxygen tubing, which is connected to the nasal prong part of the device, becomes detached (see image B above), oxygen will not be delivered to the patient. There is a risk of oxygen desaturation/hypoxaemia, which could cause life-threatening incidents.

- 2. 3. Probability of problem arising
 - 1:1,000,000 1:10,000
- 2. 4. Predicted risk to patient/users

There is a potential for a major effect on the patient health, however it has been assessed as unlikely/rare to occur.



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| 2. | 5. Further information to help characterise the problem | | | |
|----|--|--|--|--|
| | See section 2.3 above | | | |
| | | | | |
| 2. | 6. Background on Issue | | | |
| | We have received reports from two hospitals of disconnection of the tube from the nasal | | | |
| | prongs whilst in use on patients. | | | |
| | | | | |
| 2. | 7. Other information relevant to FSCA | | | |
| | N/A | | | |
| | | | | |
| | 3. Type of Action to mitigate the risk* | | | |
| | | | | |
| 3. | 1. Action To Be Taken by the User* | | | |
| | | | | |
| | ☑ Identify Device ☑ Quarantine Device ☑ Return Device ☑ Destroy Device | | | |
| | | | | |
| | | | | |
| | ☐ On-site device modification/inspection | | | |
| | | | | |
| | ☐ Follow patient management recommendations | | | |
| | | | | |
| | ☑ Take note of amendment/reinforcement of Instructions For Use (IFU) | | | |
| | ☐ Other ☐ None | | | |
| | | | | |
| | Identify and Immediately quarantine all affected lot numbers listed above and do not use | | | |
| | these devices. Please complete the Reply Form to confirm the products have been | | | |
| | · · · · · · · · · · · · · · · · · · · | | | |
| | disposed of locally or to arrange collection of the devices and a credit. If you have no | | | |
| | affected devices in stock, please confirm this using the Reply Form. Return the | | | |
| | completed Reply Form to giedriusb@intersurgical.lt (local contact e-mail address). | | | |
| | | | | |
| | Please continue to report to Intersurgical any adverse events involving this product. | | | |
| | | | | |
| | | | | |

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| | PLEASE NOTE: The potentially affected products must be removed from use as soon as replacement or alternative stock is available. We have provided these important excerpts from the IFU to help minimise the specific risk of disconnection if the products are required for use until replacement or alternative products are available. Please refer to the IFU provided for full details. PRE USE CHECK Ensure all components are undamaged and attached securely. IN USE CHECKS Patient monitoring (SaO2) must be used with this device. Ensure that nasal prongs remain inserted into the nares. Caution: 1. For use by appropriately trained personnel only. 2. Ensure that trained personnel are familiar with the contents of this instruction. 3. Always perform pre-use checks. | | | |
|----|--|--|---|--|
| 3. | | By when should the action be completed? | ongoing until no aff FSN is remaining. | eipt of this FSN and ected stock listed in this |
| 3. | 3. | Particular considerations for | r: Choose an item. | |
| | | Is follow-up of patients or re No Not applicable | eview of patients' previous resul | ts recommended? |
| 3. | 4. | Is customer Reply Required | 1? * | Yes |
| | | yes, form attached specifying | | |
| 3. | 5. | Action Being Taken by | the Manufacturer | |
| | | □ Product Removal | ☐ On-site device modification/ | inspection |
| | | □ Software upgrade | ☐ IFU or labelling change | |
| | | ○ Other | □ None | |
| | | needing to use potentially a | the Instructions For Use provid | ve products are obtained. |
| 3 | 6. | By when should the action be completed? | One month of receipt of the | e FSN |
| 3. | 7. | Is the FSN required to be co | ommunicated to the patient | Yes |
| 3 | 8. | | ovided additional information su | itable for the patient/lav |
| - | ٠. | | professional user information le | |

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| | 4. General Information* | | | |
|----|--|---|--|--|
| 4. | 1. FSN Type* | Revised FSN | | |
| 4. | For updated FSN, reference number and date of previous FSN | Superseding FSN Ref: 385183 | | |
| 4. | 3. For Updated FSN, key new infor | | | |
| | Previous advisory FSN Ref: 3 385183A. | 85183 has now been superseded by this FSN | | |
| 4. | 4. Further advice or information already expected in follow-up FSN? * | | | |
| | 5. If follow-up FSN expected, what is the further advice expected to relate to: | | | |
| 4 | | | | |
| 4 | Anticipated timescale for follow up FSN | - N/A | | |
| 4. | 7. Manufacturer information (For contact details of local representative refer to page 1 of this FSN) | | | |
| | a. Company Name | Intersurgical Ltd. | | |
| | b. Address | Crane House, Molly Millars Lane, Wokingham, Berkshire, RG41 2RZ | | |
| | c. Website address | https://www.intersurgical.com/ | | |
| 4. | 8. The Competent (Regulatory) Authority of your country has been informed about this communication to customers. * | | | |
| 4. | 9. List of attachments/appendices: | Customer Reply Form | | |
| 4. | 10. Name/Signature | Ivan Seniut, Group Quality and Regulatory Affairs Director, Intersurgical | | |
| | | <i>A</i> | | |

| Transmission of this Field Safety Notice |
|--|

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This notice needs to be passed on to all those who need to be aware within your organisation or to any organisation where the potentially affected devices have been transferred. (As appropriate)

Please transfer this notice to other organisations on which this action has an impact. (As appropriate)

Please maintain awareness on this notice and resulting action for an appropriate period to ensure effectiveness of the corrective action.

Please report all device-related incidents to the manufacturer, distributor or local representative, and the national Competent Authority if appropriate, as this provides important feedback..*

Note: Fields indicated by * are considered necessary for all FSNs. Others are optional.