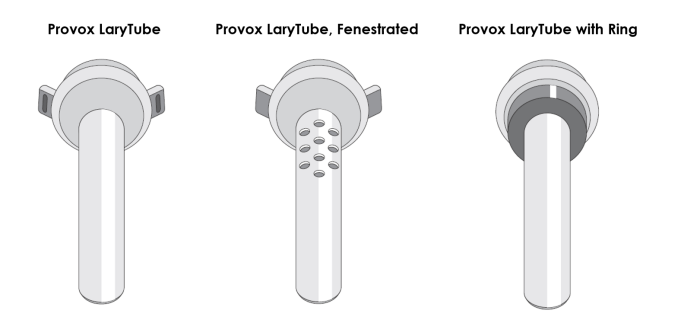


Provox[®] LaryTube[™]



Product description:

The Provox LaryTube is a tube made of medical grade silicone rubber. The purpose of the device is to create a comfortable and airtight fit between the Provox LaryTube and the tracheostoma, and to provide attachment for devices from the Provox HME System.

The Standard model and the Ring version can be fenestrated so that air can go through the voice prosthesis for voice prosthesis users.

Standard versions – made for use with or without a voice prosthesis. Can be attached with a Provox TubeHolder or Provox LaryClip.

Fenestrated versions – for voice prosthesis users. Can be attached with a Provox TubeHolder or Provox LaryClip.

Ring versions – made for use with or without a voice prosthesis.

| Atos Medical AB | SE-242 35 Hörby, Sweden | Web Site: www.atosmedical.com | Org.nr 556268-7607 |
|-----------------|-------------------------|-------------------------------|------------------------|
| Kraftgatan 8 | Tel: +46 (0) 415 198 00 | E-mail: info@atosmedical.com | VAT no. SE556268760701 |

File name:

Template ID: TMP-0260 Version: 10 Valid from: 2023/10/02



Atos Product Information

| Document ID: | PF011-01-TechInfo | Edition: | 2.0 |
|-----------------------------------|---|------------------|----------|
| Manufacturer: | Atos Medical AB Kraftgatan 8 SE-242 35 Hörby,Sweden | | |
| Classification: (EU) 93/42/EEC | IIb (Rule 5) | | |
| Intended Use: | The Provox LaryTube is a holder for devices in the Provox HME System intended for vocal and pulmonary rehabilitation after total laryngectomy. For patients with a shrinking tracheostoma it is also used to maintain the tracheostoma for breathing. The Provox LaryTube is intended for single patient use. | | |
| Use specifications: | Intended medical indication Product for rehabilitation for patients breathing the | nrough a trach | eostoma. |
| | Intended patient population Patients of any age. Cognitive ability, by a clinician judged as sufficie Manual dexterity, by a clinician judged as sufficie | | |
| | Intended usage Single patient multiple use, Prescription only. | | |
| | Intended part of the body/type of tissue applied to or interacted with Tracheostoma. | | |
| | Intended user profile The product is supposed to be handled by the patient but is also handled by physicians, trained nurses, SLPs, clinicians and caregivers. | | |
| | Intended conditions of use Environment: Home use (normal daily environme environmental restrictions regarding temperature Outpatient clinic use. Hospital use. Frequency of use: Continuous use. Replacement rate: Max use of 6 months. Replace patient, clinician or caregiver. | e, moisture etc. |). |
| Contraindications: | Provox LaryTube is not intended to be used by poare under any form of mechanical ventilation.have damaged tracheal or tracheostoma tissues | | |
| CE Mark: | Yes. Devices are CE-marked. | | |
| GMDN code: | 12292 (Laryngectomy tube) | | |
| Sterilization: | Non-sterile | | |
| Raw material: | LaryTube: Silicone Ring: Silicone with blue masterbatch | | |
| Latex information: | Not manufactured with natural rubber latex. | | |



| Biological origin: | The device is not manufactured with materials derived from human or animal source. |
|--|--|
| Handling and storage: | Store the product dry and away from sunlight at room temperature. Excursions permitted between 2 °C - 42 °C. |
| Waste handling and disposal: | Waste handling and disposal should be carried out in agreement with medical practice and applicable national laws and legislations. Used product may be a potential biohazard. |
| Hazardous components: | None. |
| Expiration date: | 3 years after manufacturing. |
| Packaging:Provox LaryTube (standard) is packed in a plastic bag of polyethylene.5 pcs of Provox XtraFlow HME are packed in a plastic bag of polyethyleThe products and instructions for use for Provox LaryTube and ProvoxXtraHME are packed in a cardboard box. | |
| | Provox LaryTube (fenestrated) is packed in a plastic bag of polyethylene. 5 pcs of Provox XtraFlow HME are packed in a plastic bag of polyethylene. 1 Provox Brush is packed in a plastic bag of polyethylene. The products and instructions for use for Provox LaryTube and Provox XtraHME and Provox Brush are packed in a cardboard box. |
| | Provox LaryTube (with ring) is packed in a plastic bag of polyethylene. 5 pcs of Provox XtraFlow HME are packed in a plastic bag of polyethylene. The products and instructions for use for Provox LaryTube and Provox XtraHME are packed in a cardboard box. |



Devices under Basic UDI-DI: 7331791-LTU-0-000-0002-3E

| REF | Name | UDI-DI |
|------|------------------------------------|----------------|
| 7601 | Provox LaryTube 8/27 | 07331791002076 |
| 7602 | Provox LaryTube 8/36 | 07331791002090 |
| 7603 | Provox LaryTube 8/55 | 07331791002113 |
| 7605 | Provox LaryTube 9/27 | 07331791002137 |
| 7606 | Provox LaryTube 9/36 | 07331791002151 |
| 7607 | Provox LaryTube 9/55 | 07331791002175 |
| 7609 | Provox LaryTube 10/27 | 07331791002199 |
| 7610 | Provox LaryTube 10/36 | 07331791002212 |
| 7611 | Provox LaryTube 10/55 | 07331791002236 |
| 7613 | Provox LaryTube 12/27 | 07331791002250 |
| 7614 | Provox LaryTube 12/36 | 07331791002274 |
| 7615 | Provox LaryTube 12/55 | 07331791002298 |
| 7624 | Provox LaryTube 8/36 with Ring | 07331791002311 |
| 7625 | Provox LaryTube 8/55 with Ring | 07331791002335 |
| 7626 | Provox LaryTube 9/36 with Ring | 07331791002359 |
| 7627 | Provox LaryTube 9/55 with Ring | 07331791002373 |
| 7628 | Provox LaryTube 10/36 with Ring | 07331791002397 |
| 7629 | Provox LaryTube 10/55 with Ring | 07331791002410 |
| 7630 | Provox LaryTube 12/36 with Ring | 07331791002434 |
| 7631 | Provox LaryTube 12/55 with Ring | 07331791002458 |
| 7637 | Provox LaryTube 8/36, Fenestrated | 07331791002472 |
| 7638 | Provox LaryTube 8/55, Fenestrated | 07331791002496 |
| 7640 | Provox LaryTube 9/36, Fenestrated | 07331791002519 |
| 7641 | Provox LaryTube 9/55, Fenestrated | 07331791002533 |
| 7643 | Provox LaryTube 10/36, Fenestrated | 07331791002557 |
| 7644 | Provox LaryTube 10/55, Fenestrated | 07331791002571 |
| 7646 | Provox LaryTube 12/36, Fenestrated | 07331791002595 |
| 7647 | Provox LaryTube 12/55, Fenestrated | 07331791002618 |

Atos Medical AB compatible products:

| Range | BASIC UDI-DI |
|-------------------------------|---------------------------|
| Provox Adhesive | 7331791-ADH-0-000-0000-CQ |
| Provox BasePlate Adaptor | 7331791-HME-A-000-0003-F5 |
| Provox Brush | 7331791-VPS-A-000-0003-RR |
| Provox FreeHands HME Cassette | 7331791-HME-0-000-0003-XJ |
| Provox LaryClip | 7331791-LTU-A-000-0001-JT |
| Provox Micron HME | 7331791-HME-0-000-0002-XF |
| Provox ShowerAid | 7331791-ADH-A-000-0000-U8 |
| Provox Swab | 7331791-GEN-A-000-0002-EC |
| Provox TubeBrush | 7331791-GEN-A-000-0001-E9 |
| Provox TubeHolder | 7331791-GEN-A-000-0000-E6 |
| Provox XtraHME | 7331791-HME-0-000-0000-X9 |

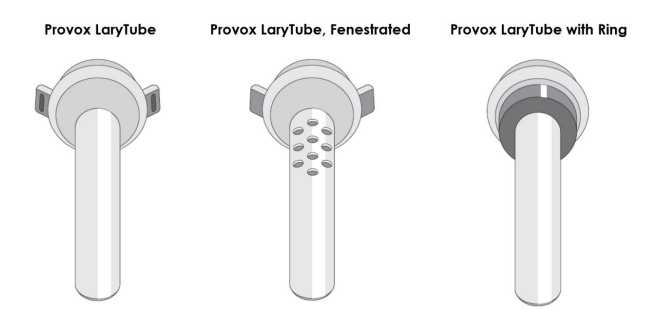
Document Approvals

Approved Date: 2023-10-24

| Task: Approval Task Verdict: Approve | ABDALM Abdallah Almashharawi, Sustaining Engineer (abdallah.almashharawi- atosmedical@coloplast.com) Issuer 23-Oct-2023 10:05:09 GMT+0000 |
|--|--|
| Task: Final Approval Verdict: Approve | ELIAND Elin Andersson, Associate Design Control & Usability Specialist (elin.andersson-atosmedical@coloplast.com) Technical / Specialist 24-Oct-2023 09:29:44 GMT+0000 |



Provox[®] LaryTube[™]



Product description:

The Provox LaryTube is a tube made of medical grade silicone rubber. The purpose of the device is to create a comfortable and airtight fit between the Provox LaryTube and the tracheostoma, and to provide attachment for devices from the Provox HME System.

The Standard model and the Ring version can be fenestrated so that air can go through the voice prosthesis for voice prosthesis users.

Standard versions – made for use with or without a voice prosthesis. Can be attached with a Provox TubeHolder or Provox LaryClip.

Fenestrated versions – for voice prosthesis users. Can be attached with a Provox TubeHolder or Provox LaryClip.

Ring versions – made for use with or without a voice prosthesis.

| Atos Medical AB | SE-242 35 Hörby, Sweden | Web Site: www.atosmedical.com | Org.nr 556268-7607 |
|-----------------|-------------------------|-------------------------------|------------------------|
| Kraftgatan 8 | Tel: +46 (0) 415 198 00 | E-mail: info@atosmedical.com | VAT no. SE556268760701 |

File name:

Template ID: TMP-0260 Version: 10 Valid from: 2023/10/02

Page 1 of 4



| Document ID: | PF011-02-TechInfo | Edition: | 2.0 |
|-----------------------------------|--|-------------------------------------|------------------------|
| Manufacturer: | Atos Medical AB Kraftgatan 8 SE-242 35 Hörby,Sweden | | |
| Classification: (EU) 93/42/EEC | llb (Rule 5) | | |
| Intended Use: | The Provox LaryTube is a holder for devices in the intended for vocal and pulmonary rehabilitation For patients with a shrinking tracheostoma it is al tracheostoma for breathing. The Provox LaryTube patient use. | after total lary so used to mair | ngectomy. ntain the |
| Use specifications: | Intended medical indication Product for rehabilitation for patients breathing t | hrough a trach | eostoma. |
| | Intended patient population Patients of any age. Cognitive ability, by a clinician judged as sufficie Manual dexterity, by a clinician judged as suffici | | |
| | Intended usage Single patient multiple use, Prescription only. | | |
| | Intended part of the body/type of tissue applied to or interacted with Tracheostoma. | | |
| | Intended user profile The product is supposed to be handled by the patient but is also handled by physicians, trained nurses, SLPs, clinicians and caregivers. | | |
| | Intended conditions of use Environment: Home use (normal daily environme environmental restrictions regarding temperature Outpatient clinic use. Hospital use. Frequency of use: Continuous use. Replacement rate: Max use of 6 months. Replace patient, clinician or caregiver. | e, moisture etc. | .). |
| Contraindications: | Provox LaryTube is not intended to be used by pare under any form of mechanical ventilation.have damaged tracheal or tracheostoma tissues | | |
| CE Mark: | Yes. Devices are CE-marked. | | |
| GMDN code: | 12292 (Laryngectomy tube) | | |
| Sterilization: | Non-sterile | | |
| Raw material: | LaryTube: Silicone Ring: Silicone with blue masterbatch | | |
| Latex information: | Not manufactured with natural rubber latex. | | |



Atos Product Information

| Biological origin: | The device is not manufactured with materials derived from human or animal source. |
|---------------------------------|--|
| Handling and storage: | Store the product dry and away from sunlight at room temperature. Excursions permitted between 2 °C - 42 °C. |
| Waste handling and disposal: | Waste handling and disposal should be carried out in agreement with medical practice and applicable national laws and legislations. Used product may be a potential biohazard. |
| Hazardous components: | None. |
| Expiration date: | 5 years after manufacturing. |
| Packaging: | Provox LaryTube is packed in a plastic bag of polyethylene. The product and instructions for use are packed in a cardboard box. |



Devices under Basic UDI-DI: 7331791-LTU-0-000-0002-3E

| REF | Name | UDI-DI |
|--------|------------------------------------|---------------|
| 7601FR | Provox LaryTube 8/27 | 7331791002083 |
| 7602FR | Provox LaryTube 8/36 | 7331791002106 |
| 7603FR | Provox LaryTube 8/55 | 7331791002120 |
| 7605FR | Provox LaryTube 9/27 | 7331791002144 |
| 7606FR | Provox LaryTube 9/36 | 7331791002168 |
| 7607FR | Provox LaryTube 9/55 | 7331791002182 |
| 7609FR | Provox LaryTube 10/27 | 7331791002205 |
| 7610FR | Provox LaryTube 10/36 | 7331791002229 |
| 7611FR | Provox LaryTube 10/55 | 7331791002243 |
| 7613FR | Provox LaryTube 12/27 | 7331791002267 |
| 7614FR | Provox LaryTube 12/36 | 7331791002281 |
| 7615FR | Provox LaryTube 12/55 | 7331791002304 |
| 7624FR | Provox LaryTube 8/36 with Ring | 7331791002328 |
| 7625FR | Provox LaryTube 8/55 with Ring | 7331791002342 |
| 7626FR | Provox LaryTube 9/36 with Ring | 7331791002366 |
| 7627FR | Provox LaryTube 9/55 with Ring | 7331791002380 |
| 7628FR | Provox LaryTube 10/36 with Ring | 7331791002403 |
| 7629FR | Provox LaryTube 10/55 with Ring | 7331791002427 |
| 7630FR | Provox LaryTube 12/36 with Ring | 7331791002441 |
| 7631FR | Provox LaryTube 12/55 with Ring | 7331791002465 |
| 7637FR | Provox LaryTube 8/36, Fenestrated | 7331791002489 |
| 7638FR | Provox LaryTube 8/55, Fenestrated | 7331791002502 |
| 7640FR | Provox LaryTube 9/36, Fenestrated | 7331791002526 |
| 7641FR | Provox LaryTube 9/55, Fenestrated | 7331791002540 |
| 7643FR | Provox LaryTube 10/36, Fenestrated | 7331791002564 |
| 7644FR | Provox LaryTube 10/55, Fenestrated | 7331791002588 |
| 7646FR | Provox LaryTube 12/36, Fenestrated | 7331791002601 |
| 7647FR | Provox LaryTube 12/55, Fenestrated | 7331791002625 |

Atos Medical AB compatible products:

| Range | BASIC UDI-DI |
|-------------------------------|---------------------------|
| Provox Adhesive | 7331791-ADH-0-000-0000-CQ |
| Provox BasePlate Adaptor | 7331791-HME-A-000-0003-F5 |
| Provox Brush | 7331791-VPS-A-000-0003-RR |
| Provox FreeHands HME Cassette | 7331791-HME-0-000-0003-XJ |
| Provox LaryClip | 7331791-LTU-A-000-0001-JT |
| Provox Micron HME | 7331791-HME-0-000-0002-XF |
| Provox ShowerAid | 7331791-ADH-A-000-0000-U8 |
| Provox Swab | 7331791-GEN-A-000-0002-EC |
| Provox TubeBrush | 7331791-GEN-A-000-0001-E9 |
| Provox TubeHolder | 7331791-GEN-A-000-0000-E6 |
| Provox XtraHME | 7331791-HME-0-000-0000-X9 |

Document Approvals

Approved Date: 2023-10-24

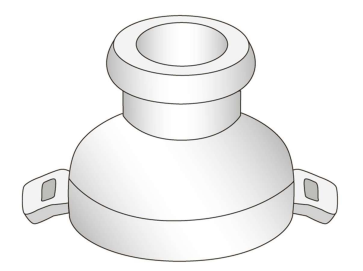
| Task: Approval Task Verdict: Approve | ABDALM Abdallah Almashharawi, Sustaining Engineer (abdallah.almashharawi- atosmedical@coloplast.com) Issuer 23-Oct-2023 10:05:08 GMT+0000 |
|--|--|
| Task: Final Approval Verdict: Approve | ELIAND Elin Andersson, Associate Design Control & Usability Specialist (elin.andersson-atosmedical@coloplast.com) Technical / Specialist 24-Oct-2023 09:27:50 GMT+0000 |

| Justification: | Function: | Electronic signature justification: | Signed: Date (yyyy-mm-dd) - Time (hh:mm): |
|----------------|-----------|-------------------------------------|---|
| Issued: | QA | Carolina Johansson - SEHRBJNC | 2022-04-12 - 14:46 |
| Reviewed: | QA | Karolina Nilsson - KARNIL | 2022-04-12 - 15:48 |
| Approved: | DD | Diana Tieger - DIATIE | 2022-04-14 - 08:06 |
| Released: | QA | Carolina Johansson - SEHRBJNC | 2022-05-19 - 15:07 |

This document has been electronically signed by the persons above.



Provox[®] LaryButton[™]



Product description:

Atos Medical AB

Kraftgatan 8

Provox LaryButton is delivered single packed, non-sterile, ready for use. The goal is to create a self-retaining, comfortable and airtight fit between the Provox LaryButton and the tracheostoma.

| File name: PF031-01-Techinfo | |
|---|---|
| Template ID: TMP-0260 Deocumental Number/02/2V-0545293 Status: Effective Version: 1.0 | 0 |

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E-mail:

Web Site: www.atosmedical.com

info@atosmedical.com

Org.nr 556268-7607

VAT no. SE556268760701



| Document ID: | PF031-01-TechInfo | Edition: | 06 |
|---------------------------------------|---|--|--|
| Manufacturer: | Atos Medical AB Kraftgatan 8 SE-242 35 Hörby,Sweden | | |
| Classification: (EU) MDD 93/42/EEC | llb (2.1 Rule 5) | | |
| Intended Use: | The Provox LaryButton is a self-retaining holder for HME System intended for vocal and pulmonary re laryngectomy. For patients with a shrinking tracheostomas it is a tracheostoma for beathing. The Provox LaryButton is intended for single patie | ehabilitation a Iso used to ma | fter total |
| Use specifications: | Intended medical indication Product for rehabilitation for patients breathing to Intended patient population Patients of any age. Cognitive ability, by a clinician judged as sufficients Manual dexterity, by a clinician judged as sufficients Manual dexterity, by a clinician judged as sufficients Not intended for patients with mechanical ventilents Not intended for patients with a low tidal volume Intended usage Provox LaryButton is a single patient use device provided the body/type of tissue applied Tracheostoma. Intended user profile The product is supposed to be handled by the prover by physicians, trained nurses, SLPs, clinicians and Intended conditions of use Environment: Home use (normal daily environment environmental restrictions regarding temperatures Outpatient clinic use. Hospital use. Frequency of use: Continuous use. Replacement rate: Max usage for 6 months. Replacement rate: Max usage for 6 months. Replacement, clinician or caregiver. | ent. ent. ation. e. prescribed by c to or interacte atient but is als caregivers. nt without any e, moisture etc. | a clinician. d with so handled or .). |
| Contraindications: | Provox LaryButton is not intended to be used by form of mechanical ventilation or have damage | | |
| CE Mark: | Yes. Devices are CE-marked. | | |
| GMDN code: | 14093 (Tracheostoma button) | | |
| Sterilization: | Non-sterile | | |
| Raw material: | Silicone | | |
| Latex information: | Not manufactured with natural rubber latex | | |
| Biological origin: | The device is not manufactured with materials d animal source. | erived from hu | man or |
| Handling and storage: | Store the product dry and away from sunlight at Excursions permitted between 2°C - 42°C. | room temperc | iture. |



| Waste handling and disposal: | Waste handling and disposal should be carried out in agreement with medical practice and applicable national laws and legislations. Used product may be a potential biohazard. |
|---------------------------------|--|
| Hazardous components: | None |
| Expiration date: | 5 years after manufacturing. |
| Packaging: | Provox LaryButton is packed in a plastic bag of polyethylene. The product and instructions for use are packed in a cardboard box. |

Devices under Basic UDI-DI: 7331791-LTU-0-000-0000-38

| REF | Name | UDI-DI |
|------|-------------------------|----------------|
| 7671 | Provox LaryButton 12/8 | 07331791002694 |
| 7672 | Provox LaryButton 14/8 | 07331791002700 |
| 7673 | Provox LaryButton 16/8 | 07331791002717 |
| 7674 | Provox LaryButton 18/8 | 07331791002724 |
| 7685 | Provox LaryButton 12/18 | 07331791002731 |
| 7686 | Provox LaryButton 14/18 | 07331791002748 |
| 7687 | Provox LaryButton 16/18 | 07331791002755 |
| 7688 | Provox LaryButton 18/18 | 07331791002762 |

Atos Medical AB compatible products:

| Range | BASIC UDI-DI |
|-------------------------------|---------------------------|
| Provox BasePlate Adaptor | 7331791-HME-A-000-0003-F5 |
| Provox FreeHands HME Cassette | 7331791-HME-0-000-0003-XJ |
| Provox LaryClip | 7331791-LTU-A-000-0001-JT |
| Provox Micron HME | 7331791-HME-0-000-0002-XF |
| Provox ShowerAid | 7331791-ADH-A-000-0000-U8 |
| Provox Swab | 7331791-GEN-A-000-0002-EC |
| Provox TubeBrush | 7331791-GEN-A-000-0001-E9 |
| Provox TubeHolder | 7331791-GEN-A-000-0000-E6 |
| Provox XtraHME | 7331791-HME-0-000-0000-X9 |



| Justification: | Function: | Electronic signature justification: | Signed: Date (yyyy-mm-dd) - Time (hh:mm): |
|----------------|-----------|-------------------------------------|---|
| Issued: | QA | Niki Svensson - NIKSVE | 2022-12-19 - 11:36 |
| Reviewed: | QA | Sofia Thomasson - SOFTHO | 2022-12-19 - 11:42 |
| Approved: | QA | Elin Andersson - ELIAND | 2022-12-22 - 15:47 |
| Released: | QA | Niki Svensson - NIKSVE | 2023-03-16 - 13:32 |

This document has been electronically signed by the persons above.



Product description:

The Sizer Kit is a box which contains samples, (Sizers.) of commercially available Provox LaryButtons. The sizes of these Sizers and actual Provox LaryButtons are the same and are indicated on the products themselves and in the bottom of the outer storage box. Each Sizer in the Sizer Kit is stored in an individual removable polypropylene box. This makes it possible for the prescribing specialist to remove the individual storage boxes with the Sizers from the outer storage box individually. This allows for hygienic handling of both the Sizers and the storage boxes. After each sizing session, the Sizer(s) with its individual storage box(es) must be cleaned, disinfected, dried and steam sterilized according to the accompanying Instructions for cleaning and sterilization. The outer storage box must also be cleaned if contaminated. The Sizer LaryButtons and their individual removable storage boxes are thereafter put back at the appropriate position as indicated in the bottom of the outer storage box

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| Document ID: | PF032-01-TechInfo | Edition: | 04 | |
|------------------------------------|---|---|--|--------------------------|
| Manufacturer: | Atos Medical AB Kraftgatan 8 SE-242 35 Hörby,Sweden | | | |
| Classification: (MDD 93/42/EEC) | lla (2.1 Rule 5) | | | |
| Intended Use: | The Provox® LaryButton Sizer Kit is intended for clinician to determine the size(s) of LaryButton the patient. The Sizer Kit should be used only has read the LaryButton Manual. A copy of the Sizer Kit. It can also be viewed on the Internet The Sizer LaryButtons are intended for the size correct size(s) have been determined a new prescribed to the patient for actual use. | on that should be p by a prescribing c that manual come at at www.atosmec ng procedure only | rescribed to linician who s with the dical.com. r. After the | |
| Use specifications: | Intended medical condition Laryngectomized patient. | | | |
| | Intended patient population Gender: Male and female. Age: Typical average age for a laryngectom Intended usage | ny is 65 years. | | Release date: 2023-03-16 |
| | The Sizer LaryButtons are intended for the sizi | | | ase da |
| | Intended part of the body/type of tissue app Neck | blied to or interacte | d with | 4 Rele |
| | Intended user profile Prescribing clinician. | | | Edition: 04 |
| | Intended conditions of use Only to be used in clinical environment. | | | 0038474 |
| Contraindications: | The Sizer Kit in itself does not have specific co Provox LaryButton, or use it only with special tissue problems such as damaged mucous n formation, and vulnerability with a higher ter LaryButton may be contraindicated for patie undergoing anticoagulant treatment. | care, in cases of tr nembrane, granulo ndency to bleed. T | acheostoma ation tissue he Provox | Document No: 1000 |
| CE Mark: | Yes. Device is CE-marked. | | | |
| GMDN code: | 14093 (Tracheostomy button) | | | 0 |
| Sterilization: | Non-sterile, steam sterilizable | | | SG |
| Raw material: | Silicone, polypropylene | | | S |
| Latex information: | Not manufactured with natural rubber latex | | | Π |
| Biological origin: | The device is not manufactured with materic animal source. | als derived from hu | iman or | Selea |
| | | | | Ň |

File name: PF032-01-Techinfo.docx

Document Number: VV-0543464 Status: Effective Version: 1.0 Name: PF032-01-Techinfo



| Handling and storage: | Store the product dry and away from sunlight at room temperature. Excursions permitted between 2°C - 42°C. |
|---------------------------------|---|
| Waste handling and disposal: | Waste handling and disposal should be carried out in agreement with medical practice and applicable national laws and legislations. Used product may be a potential biohazard. |
| Hazardous components: | None |
| Expiration date: | 5 years after manufacturing. |
| Packaging: | Provox LaryButton Sizer Kit is single packed in a tamper-proof plastic bag made of polypropylene together with one IFU for the product, one IFU for Provox LaryButton and one IFU for cleaning and sterilization. |



Devices under Basic UDI-DI: 7331791-LTU-0-000-0001-3B

| REF | Name | UDI-DI |
|------|-----------------------------|---------------|
| 7690 | Provox LaryButton Sizer Kit | 7331791002779 |

Atos Medical AB compatible products:

| Range | BASIC UDI-DI |
|-------|--------------|
| N/A | N/A |

| Justification: | Function: | Electronic signature justification: | Signed: Date (yyyy-mm-dd) - Time (hh:mm): |
|----------------|-----------|-------------------------------------|---|
| Issued: | DD | Pontus Eklund - X-PONEKL | 2020-04-21 - 08:24 |
| Reviewed: | DD | Jon Berg - JONBER | 2020-04-21 - 08:58 |
| Approved: | DD | Fredrik Calais - FRECAL | 2020-04-21 - 16:07 |
| Released: | DD | Pontus Eklund - X-PONEKL | 2020-10-28 - 16:39 |

This document has been electronically signed by the persons above.



Provox® Fenestration Punch



Product description:

The Fenestration Punch is used for making small fenestrations in a Provox LaryTube at desired locations.

The Fenestration Punch is made of polypropylene, stainless steel and silicone and is used for making small fenestrations in a Provox LaryTube. This is done when the Provox LaryTube is intended to be used in combination with a voice prosthesis.

 Atos Medical AB
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 Web Site: www.atosmedical.com

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 Tel: +46 (0) 415 198 00
 E-mail: info@atosmedical.com

VAT no. SE556268760701

Org.nr 556268-7607

File name: PF037-01-TECHINFO Fenestration Punch.docx Template ID: TMP-0260 Documental Number: 0543466 Status: Effective Version: 1.0 Name: PF037-01-TECHINFO Fenestration Punch

Page 1 of 3



| Document ID: | PF037-01-TechInfo | Edition: | 06 |
|----------------------------------|---|------------------|---------------|
| Manufacturer: | Atos Medical AB Kraftgatan 8, P.O. Box 183, SE-242 22 Hörby, Sweden | | |
| Classification: (EU) 2017/745 | Class I (1.1 Rule 1) | | |
| Intended Use: | The Fenestration Punch is used for making small fer LaryTube at desired locations. | nestrations in a | Provox |
| CE Mark: | Yes, the devices are CE marked. | | |
| GMDN code: | 38792 (Basic tracheostomy tube, reusable) | | |
| Sterilization: | Non-sterile | | |
| Raw material: | Stainless Steel, Plastic, Silicone | | |
| Latex information: | Not manufactured with natural rubber latex | | |
| Biological origin: | The device is not manufactured with materials der source. | ved from hum | an or animal |
| Handling and storage: | Store the product dry and away from sunlight at roopermitted between 2°C - 42°C. | om temperatur | e. Excursions |
| Waste handling and disposal: | Waste handling and disposal should be carried ou medical practice and applicable national laws an product may be a potential biohazard. | • | |
| Hazardous components: | None | | |
| Expiration date: | 5 years after manufacturing. | | |
| Packaging: | The Fenestration Punch is single-packed in a plastic bag | | |



Devices under Basic UDI-DI: 7331791-LTU-A-000-0000-JQ

| REF | Name | UDI-DI |
|------|--------------------------|----------------|
| 7654 | Provox FenestrationPunch | 07331791002632 |

Atos Medical AB Compatible products:

| Range | BASIC UDI-DI |
|-----------------|---------------------------|
| Provox LaryTube | 7331791-LTU-0-000-0002-3E |

| Justification: | Function: | Electronic signature justification: | Signed: Date (yyyy-mm-dd) - Time (hh:mm): |
|----------------|-----------|-------------------------------------|---|
| Issued: | DD | Pontus Eklund - X-PONEKL | 2020-04-20 - 13:35 |
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| Released: | DD | Pontus Eklund - X-PONEKL | 2020-10-28 - 16:35 |

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Provox® TubeBrush



Product description:

The Provox TubeBrush is used for cleaning of the Provox LaryTube and Provox LaryButton ex situ. The Provox TubeBrush is packed 6 pieces in a plastic bag. It is available in two different models with outer diameter 8 mm or 12 mm.

Atos Medical ABSE-242 22 Hörby, SwedenWeb Site:www.atosmedical.comKraftgatan 8, P.O Box 183Tel: +46 (0) 415 198 00E-mail:info@atosmedical.com

Org.nr 556268-7607

VAT no. SE556268760701

File name: PF052-01-TECHINFO Provox TubeBrush.docx Template ID: TMP-0260 Documental Number 42V-0543145 Status: Effective Version: 1.0 Name: PF052-01-TECHINFO Provox TubeBrush



| Document ID: | PF052-01-TechInfo | Edition: | 09 |
|----------------------------------|---|------------------|----------------|
| Manufacturer: | Atos Medical AB Kraftgatan 8, P.O. Box 183, SE-242 22 Hörby, Sweden | | |
| Classification: (EU) 2017/745 | Class I, Rule 1 | | |
| Intended Use: | The Provox TubeBrush is used for cleaning of the Pro LaryButton ex situ. | ovox LaryTube | and Provox |
| CE Mark: | Yes, the devices are CE marked. | | |
| GMDN code: | 34883 (Airway device, cleaning brush, noninvasive |). | |
| Sterilization: | Non-Sterile | | |
| Raw material: | ABS, Stainless Steel, PBT and Cotton. | | |
| Latex information: | Not manufactured with natural rubber latex | | |
| Biological origin: | The device is not manufactured with materials der source. | ved from hum | an or animal |
| Handling and storage: | Store the product dry and away from sunlight at roopermitted between 2°C - 42°C. | om temperatu | re. Excursions |
| Waste handling and disposal: | Waste handling and disposal should be carried ou medical practice and applicable national laws an product may be a potential biohazard. | | |
| Hazardous components: | None | | |
| Expiration date: | 3 years after manufacturing. | | |
| Packaging: | 6 pieces Provox TubeBrush are packed in a tampe with Instructions for Use. | rproof plastic b | ag together |



Devices under Basic UDI-DI: 7331791-GEN-A-000-0001-E9

| REF | Name | UDI-DI |
|------|------------------------|---------------|
| 7660 | Provox TubeBrush 8 mm | 7331791002656 |
| 7661 | Provox TubeBrush 12 mm | 7331791002663 |

Atos Medical AB Compatible products:

| Range | BASIC UDI-DI |
|-----------------------------|---------------------------|
| Provox LaryTube | 7331791-LTU-0-000-0002-3E |
| Provox LaryButton | 7331791-LTU-0-000-0000-38 |
| Provox LaryButton Sizer Kit | 7331791-LTU-0-000-0001-3B |

| Justification: | Function: | Electronic signature justification: | Signed: Date (yyyy-mm-dd) - Time (hh:mm): |
|----------------|-----------|-------------------------------------|---|
| Issued: | QA | Sara Dahl - X-SARDAH | 2021-11-11 - 18:58 |
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| Approved: | DD | Diana Tieger - DIATIE | 2021-11-16 - 16:11 |
| Released: | QA | Sara Dahl - X-SARDAH | 2021-12-10 - 09:53 |

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Provox® TubeHolder



Product description:

The Provox TubeHolder has been developed for use with the Provox LaryTube and Provox LaryButton. The integrated clip connectors allow for optimal fit to the wings of the Provox LaryTube and LaryButton, which reduces the physical stress on the soft silicone material.

| Atos Medical AB | SE-242 35 Hörby, Sweden | Web Site: www.atosmedical.com | Org.nr 556268-7607 |
|-----------------|-------------------------|-------------------------------|------------------------|
| Kraftgatan 8 | Tel: +46 (0) 415 198 00 | E-mail: info@atosmedical.com | VAT no. SE556268760701 |
| | . , | | |

File name: PF053-01-TECHINFO Provox TubeHolder.docx Template ID: TMP-0260 Documental Number 2020-0543442 Status: Effective Version: 1.0 Name: PF053-01-TECHINFO Provox TubeHolder



| Document ID: | PF053-01-TechInfo | Edition: | 07 |
|----------------------------------|---|--|--------------------|
| Manufacturer: | Atos Medical AB Kraftgatan 8 SE-242 35 Hörby, Sweden | | |
| Classification: (EU) 2017/745 | Class I (Rule 1) | | |
| Intended Use: | The Provox TubeHolder is used for extra support for Provox LaryTube. It goes around the neck of the attached to the "ears" of the LaryTube/LaryButto adjustable in length using a Velcro® connection the band to suitable length. | user and the e | nds are Ider is |
| Use specifications: | Intended medical indication: Patients breathing through a tracheostoma. Intended patient population: Patients of any age. Cognitive ability, by a clinician judged as sufficient Manual dexterity, by a clinician judged as sufficient Intended usage: Single use. Intended part of the body/type of tissue applied The device will contact intact skin on the neck. Intended user profile: The product is supposed to be handled by the p by physicians, trained nurses, SLPs, clinicians and Intended conditions of use: Environment: Home use (normal daily environme environmental restrictions regarding temperature use. Frequency of use: Continuous use. | ent. to or interacte atient but is als caregivers. nts without any | o handled / |
| Contraindications: | None. | | |
| CE Mark: | Yes. Devices are CE-marked. | | |
| GMDN code: | 63438 (Tracheostomy tube neck holder, single-us | e) | |
| Sterilization: | Non-Sterile | | |
| Raw material: | Tricot textile, Polyurethane (PUR) foam, Polyamid | e (PA). | |
| Latex information: | Not manufactured with natural rubber latex | | |
| Biological origin: | The device is not manufactured with materials de animal source. | erived from hu | man or |
| Handling and storage: | Store the product dry and away from sunlight at Excursions permitted between 2°C - 42°C. | room temperc | iture. |
| Waste handling and disposal: | Waste handling and disposal should be carried a medical practice and applicable national laws a product may be a potential biohazard. | - | |



| Hazardous components: | None. |
|--------------------------|---|
| Expiration date: | 5 years after manufacturing. |
| Packaging: | Single packed together with IFU in a plastic bag. |

Devices under Basic UDI-DI: 7331791-GEN-A-000-0000-E6

| REF | Name | UDI-DI |
|------|-------------------|----------------|
| 7668 | Provox TubeHolder | 07331791002670 |

Atos Medical AB compatible products:

| Range | BASIC UDI-DI |
|-----------------------------|---------------------------|
| Provox LaryTube | 7331791-LTU-0-000-0002-3E |
| Provox LaryButton | 7331791-LTU-0-000-0000-38 |
| Provox LaryButton Sizer Kit | 7331791-LTU-0-000-0001-3B |
| Provox Life LaryTube | 7331791-LTU-0-000-0004-3L |
| Provox Life LaryButton | 7331791-LTU-0-000-0005-3P |

| Justification: | Function: | Electronic signature justification: | Signed: Date (yyyy-mm-dd) - Time (hh:mm): |
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| Issued: | QA | Abdallah Almashharawi - ABDALM | 2022-07-25 - 08:36 |
| Reviewed: | QA | Karolina Nilsson - KARNIL | 2022-07-25 - 09:15 |
| Approved: | DD | Peter Sundsten - PETSUN | 2022-07-27 - 08:09 |
| Released: | QA | Abdallah Almashharawi - ABDALM | 2022-07-29 - 09:09 |

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Provox[®] LaryClip[™]



Product description:

The Provox LaryClip consists of a square adhesive base and a hook-and-loop clip that allows for optimal fit to the wings of the Provox LaryButton and LaryTube. When the adhesive Base is attached to the skin at both sides of the stoma and is eventually removed due to loss of its stickiness, the Clip part can be removed and re-attached as needed.

Atos Medical AB Kraftgatan 8 SE-242 35 Hörby, Sweden Tel: +46 (0) 415 198 00 Web Site: www.atosmedical.com E-mail: info@atosmedical.com

VAT no. SE556268760701

Org.nr 556268-7607

File name: PF061-01-TECHINFO Template ID: TMP-0260 Documental Number: 01/2V-0543443 Status: Effective Version: 1.0 Name: PF061-01-TECHINFO



| Document ID: | PF061-01-TechInfo | Edition: | 08 | |
|----------------------------------|---|--|---------------|--|
| Manufacturer: | Atos Medical AB Kraftgatan 8 SE-242 35 Hörby,Sweden | | | |
| Classification: (EU) 2017/745 | Class I (1.1, Rule 1) | | | |
| Intended Use: | The Provox LaryClip is used for extra support for LaryButton and LaryTube. The product consists of two parts, one that is attached to the patients' skin on each side of the stoma and the other part is attached to the LaryButton or the LaryTube. The two parts are then connected by Velcro. | | atients' skin | |
| Use specifications: | : Intended medical indication: Product for rehabilitation for patients breathing through a tracheostoma. | | | |
| | Intended patient population: Patients of any age. Cognitive ability, by a clinician judged as sufficient. Manual dexterity, by a clinician judged as sufficient. | | | |
| | Intended usage: Single use, Over-the-counter | | | |
| | Intended part of the body/type of tissue applied to or interacted with: The device will contact intact skin (neck). | | | |
| | Intended user profile: The product is supposed to be handled by the patient but is also handled by physicians, trained nurses, SLPs, clinicians and caregivers. | | | |
| | environmental restrictions regarding temperature Outpatient clinic use. Hospital use. Frequency of use: Continuous use. | ment: Home use (normal daily environment without any or mental restrictions regarding temperature, moisture etc.). tient clinic use. Hospital use. | | |
| Contraindications: | There are no known contraindications. | | | |
| CE Mark: | Yes. Devices are CE-marked. | | | |
| GMDN code: | 35752 (Tracheostomy tube neck holder, reusable | .) | | |
| Sterilization: | Non-sterile | | | |
| Raw material: | LaryClip Base: Polyethylene (PE), Acrylic Adhesiv LaryClip: Knitted fabric, Polyamide (PA) | e, velcro | | |
| Latex information: | Not manufactured with natural rubber latex. | | | |
| Biological origin: | The device is not manufactured with materials de animal source. | erived from hur | man or | |
| Handling and storage: | Store the product dry and away from sunlight at Excursions permitted between 2°C - 42°C. | room tempera | ture. | |
| | | | | |



| Waste handling and disposal: | Waste handling and disposal should be carried out in agreement with medical practice and applicable national laws and legislations. Used product may be a potential biohazard. |
|---------------------------------|--|
| Hazardous components: | None. |
| Expiration date: | 3 years after manufacturing. |
| Packaging: | One package consists of 8 pcs of LaryClip and 40 pcs of LaryClip Base. They are packed together with instruction for use in a cardboard box. |

Devices under Basic UDI-DI: 7331791-LTU-A-000-0001-JT

| REF | Name | UDI-DI |
|------|-----------------|----------------|
| 7669 | Provox LaryClip | 07331791002687 |

Atos Medical AB compatible products:

| Range | BASIC UDI-DI |
|-----------------------------|---------------------------|
| Provox LaryTube | 7331791-LTU-0-000-0002-3E |
| Provox LaryButton | 7331791-LTU-0-000-0000-38 |
| Provox LaryButton Sizer Kit | 7331791-LTU-0-000-0001-3B |
| Provox Vega Plug 17 | 7331791-VPS-A-000-0004-RU |
| Provox Vega Plug 20 | 7331791-VPS-A-000-0004-RU |
| Provox Vega Plug 22.5 | 7331791-VPS-A-000-0004-RU |



Provox® LaryTube™ Sizer Kit



Product description:

The Sizer Kit is a box which contains samples ("sizers") of a variety of commercially available Provox LaryTubes. The sizes of these Sizers and actual Provox LaryTubes are the same. The size is indicated on the products and both diameter and length are indicated on the chart inside the box. Each sizer in the Sizer Kit is stored in an individual removable polypropylene box. This makes it possible for the prescribing specialist to remove the individual storage boxes with the Sizers from the outer storage box individually. This allows for hygienic handling of both the Sizer(s) and the storage box. After each sizing session, the Sizer(s) with its individual storage box must be cleaned, disinfected, dried and steam sterilized according to the accompanying "instructions for cleaning and sterilization".

File name: Template ID: TMP-0260 Version: 10 Valid from: 2023/10/02



Atos Product Information

| Document ID: | PF062-01-TechInfo | Edition: | 2.0 |
|---|---|--|---------------------------------|
| Manufacturer: | Atos Medical AB Kraftgatan 8 SE-242 35 Hörby,Sweden | | |
| Classification: (EU) 93/42/EEC | lla (Rule 5) | | |
| Intended Use: | The Provox LaryTube Sizer Kit is intended for use by the prescribing specialist to determine the size(s) of LaryTube that should be prescribed to the patient. The Sizer Kit should be used only by a prescribing specialist who has read the LaryTube Manual. A copy of that manual comes with the Sizer Kit. It can also be viewed on the Internet at www.atosmedical.com. The Sizer LaryTubes are intended for the sizing procedure only. After the correct size(s) have been determined, new LaryTube(s) shall be given to the patient for use. | | |
| Use specifications: | Intended medical condition Laryngectomized patient. | | |
| | Intended patient population | | |
| | Gender: Male and female. | | |
| | Age: Typical average age for a laryngectomy is | 65 years. | |
| | Intended usage The Sizer LaryTubes are intended for the sizing procedure only. | | |
| | Intended part of the body/type of tissue applied to or interacted with Neck | | |
| | | | |
| | | | |
| | Neck Intended user profile | | |
| Contraindications: | Neck Intended user profile Prescribing clinician. Intended conditions of use | ontraindication | s. The Provox |
| Contraindications: CE Mark: | Neck Intended user profile Prescribing clinician. Intended conditions of use Only to be used in clinical environment. The Sizer Kit in itself does not have specific co LaryTubes contained in the LaryTube Sizer Kit ar | ontraindication | s. The Provox |
| | Neck Intended user profile Prescribing clinician. Intended conditions of use Only to be used in clinical environment. The Sizer Kit in itself does not have specific co LaryTubes contained in the LaryTube Sizer Kit ar requiring mechanical ventilation. | ontraindication | s. The Provox |
| CE Mark: | Neck Intended user profile Prescribing clinician. Intended conditions of use Only to be used in clinical environment. The Sizer Kit in itself does not have specific con LaryTubes contained in the LaryTube Sizer Kit ar requiring mechanical ventilation. Yes. Device is CE-marked. | ontraindication | s. The Provox |
| CE Mark: GMDN code: | Neck Intended user profile Prescribing clinician. Intended conditions of use Only to be used in clinical environment. The Sizer Kit in itself does not have specific co LaryTubes contained in the LaryTube Sizer Kit ar requiring mechanical ventilation. Yes. Device is CE-marked. 12292 (Laryngectomy tube) | ontraindication | s. The Provox |
| CE Mark: GMDN code: Sterilization: | Neck Intended user profile Prescribing clinician. Intended conditions of use Only to be used in clinical environment. The Sizer Kit in itself does not have specific co LaryTubes contained in the LaryTube Sizer Kit ar requiring mechanical ventilation. Yes. Device is CE-marked. 12292 (Laryngectomy tube) Non-sterile, steam sterilizable. | ontraindication | s. The Provox |
| CE Mark: GMDN code: Sterilization: Raw material: | Neck Intended user profile Prescribing clinician. Intended conditions of use Only to be used in clinical environment. The Sizer Kit in itself does not have specific contaryTubes contained in the LaryTube Sizer Kit arrequiring mechanical ventilation. Yes. Device is CE-marked. 12292 (Laryngectomy tube) Non-sterile, steam sterilizable. Silicone, Polypropylene. | ontraindication: e not intended | s. The Provox d for patients |
| CE Mark: GMDN code: Sterilization: Raw material: Latex information: | Neck Intended user profile Prescribing clinician. Intended conditions of use Only to be used in clinical environment. The Sizer Kit in itself does not have specific con LaryTubes contained in the LaryTube Sizer Kit and requiring mechanical ventilation. Yes. Device is CE-marked. 12292 (Laryngectomy tube) Non-sterile, steam sterilizable. Silicone, Polypropylene. Not manufactured with natural rubber latex. The device is not manufactured with materials d | entraindication: The not intended erived from hu | s. The Provox d for patients |

File name:



| Waste handling and disposal: | Waste handling and disposal should be carried out in agreement with medical practice and applicable national laws and legislations. Used product may be a potential biohazard. |
|---------------------------------|---|
| Hazardous components: | None. |
| Expiration date: | 5 years after manufacturing. |
| Packaging: | Provox LaryTube Sizer Kit is single packed in a tamper-proof plastic bag together with a manual for the product, instructions for sterilization and a manual for the Provox LaryTube. |



Devices under Basic UDI-DI: 7331791-LTU-0-000-0003-3H

| REF | Name | UDI-DI |
|------|---------------------------|----------------|
| 7648 | Provox LaryTube Sizer Kit | 07331791005329 |

Atos Medical AB compatible products:

| Range | BASIC UDI-DI |
|-------|--------------|
| N/A | N/A |

Document Approvals

Approved Date: 2023-10-23

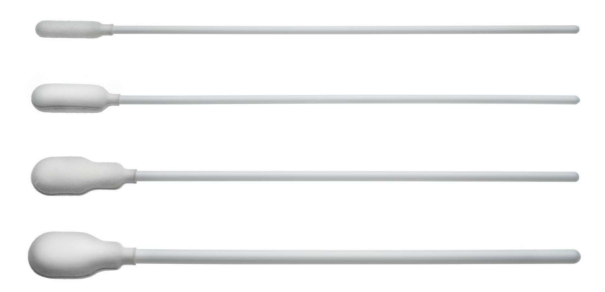
| Task: Approval Task Verdict: Approve | ABDALM Abdallah Almashharawi, Sustaining Engineer (abdallah.almashharawi- atosmedical@coloplast.com) Issuer 16-Oct-2023 07:34:12 GMT+0000 |
|--|--|
| Task: Final Approval Verdict: Approve | ELIAND Elin Andersson, Associate Design Control & Usability Specialist (elin.andersson-atosmedical@coloplast.com) Technical / Specialist 23-Oct-2023 09:09:38 GMT+0000 |

| Justification: | Function: | Electronic signature justification: | Signed: Date (yyyy-mm-dd) - Time (hh:mm): |
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| Released: | QA | Sara Dahl - X-SARDAH | 2021-12-10 - 09:51 |

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Provox[®] Swab



Product description:

The Provox Swab is a foam attached to a polymer stick handle.

| Atos Medical AB | SE-242 35 Hörby, Sweden | Web Site: www.atosmedical.com | Org.nr 556268-7607 |
|-----------------|-------------------------|-------------------------------|-------------------------|
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| | | | |



| Document ID: | PF085-01-TechInfo | Edition: | 06 |
|----------------------------------|--|----------------|----------|
| Manufacturer: | Atos Medical AB Kraftgatan 8 SE-242 35 Hörby, Sweden | | |
| Classification: (EU) 2017/745 | Class I, Rule 1 | | |
| Intended Use: | Provox Swab is a single use swab for ex-situ cleaning of Provox LaryTube, Provox LaryButton and tracheostomy inner tubes. | | |
| Use specifications: | Intended medical indication: Product for laryngectomized or tracheostomized patients, and/or their caregivers, using Provox LaryTube, Provox LaryButton or double lumen tracheostomy tube, that requires regular cleaning ex-situ. Intended patient population: Male and female, laryngectomized or tracheostomized patients. Intended usage: Single patient use, swabs should be discarded after use. Intended part of the body/type of tissue applied to or interacted with: N/A, cleaning will be performed ex-situ. Intended user profile: Patient, clinician, caregiver. Intended conditions of use: Normal daily environment without any hygienic or environmental restrictions regarding temperature, moisture etc. | | |
| Contraindications: | No identified or known contraindications. | | |
| CE Mark: | Yes. Devices are CE-marked. | | |
| GMDN code: | 62956 (Airway device cleaning utensil, noninvasi | ve, single-use |) |
| Sterilization: | Non-Sterile | | |
| Raw material: | Polypropylene (stick handle) and Polyurethane, reticulated foam (foam mitt). | | am (foam |
| Latex information: | Not manufactured with natural rubber latex. | | |
| Biological origin: | The device is not manufactured with materials derived from human or animal source. | | |
| Handling and storage: | Store the product dry and away from sunlight at room temperature. Excursions permitted between 2°C - 42°C. | | |
| Waste handling and disposal: | Waste handling and disposal should be carried out in agreement with medical practice and applicable national laws and legislations. Used product may be a potential biohazard. | | |
| Hazardous components: | None | | |
| Expiration date: | 3 years after manufacturing. | | |



Packaging:

50 pcs per package. Devices are packed in plastic bags made of polyethylene and packed together in a cardboard box with printed instructions for use.

Swab Medium is also available as 10pcs, packed in plastic bags with instructions for use printed on the label.

Devices under Basic UDI-DI: 7331791-GEN-A-000-0002-EC

| REF | Name | UDI-DI |
|------|--------------------------|----------------|
| 8250 | Provox Swabs Small | 07331791011412 |
| 8251 | Provox Swab Medium | 07331791011429 |
| 8252 | Provox Swab Large | 07331791011436 |
| 8258 | Provox Swab XtraLarge | 07331791012730 |
| 8083 | Provox Swab Medium 10pcs | 07331791016028 |

Atos Medical AB compatible products:

| Range | BASIC UDI-DI |
|------------------------|---------------------------|
| Provox LaryTube | 7331791-LTU-0-000-0002-3E |
| Provox LaryButton | 7331791-LTU-0-000-0000-38 |
| Provox Life LaryTube | 7331791-LTU-0-000-0004-3L |
| Provox Life LaryButton | 7331791-LTU-0-000-0005-3P |