



Circular No. ١٢١ / 2024

24 -02-1446 H
29 -08-2024

نتقدم بثقة
Moving Forward
with Confidence



Recall of Nasopharyngeal Airway, Double Swivel Connector, 15mm Y Piece, Portex™ Orator Speaking Valve, Thermovent™ 1200 from Smiths Medical.

Source	SFDA- Saudi Food & Drug Authority. https://ade.sfda.gov.sa/Fsca/PublishDetails/57
Product	Nasopharyngeal Airway, Double Swivel Connector, 15mm Y Piece, Portex™ Orator Speaking Valve, Thermovent™ 1200.
Manufacturer	Smiths Medical.
Local agent	Muscat Pharmacy & Stores LLC.
The affected products	Refer to Table 1 in the attachment for affected lots (manufactured between 20 January 2021 and 27 August 2021).
Reason	There is uncertainty in the seal integrity of the sterile packaging of sterilized above-mentioned products.
Action	1. Discontinue use and discard all affected products following your institution's process for discarding. If discarding is not immediately possible at your facility, then the product should be quarantined until disposal. 2. Contact the local agent for remedial action.
comments	Healthcare professionals are encouraged to report any adverse events Suspected to be associated with the above device or any other medical device to Department of Medical Device Control through the E-mail: vigilance-md@moh.gov.om

Ph. Ahmed Al Harbi
Acting Director General





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To:

THE DIRECTOR GENERAL OF HEALTH SERVICES IN ALL GOVERNORATES

Commanding Officer, Armed Forces Hospital (Al Khoudh & Salalah)

Director General of Engineering Affairs, MOH

Director General of Royal Hospital

Director General of Khoula Hospital

Director General of Medical Supplies (MOH)

Director General of Pvt. Health Est. Affairs (to kindly arrange distribution to all Pvt. Hospitals)

Hospital Director (Al Nahda Hospital)

Hospital Director (Al Massara Hospital)

The Head of Medical Services in SQU Hospital

The Head of Medical Services in Royal Oman Police

The Head of Medical Services in Ministry of Defence

The Head of Medical Services in The Diwan

The Head of Medical Services in The Sultan's Special Force

The Head of Medical Services in Internal Security Services

The Head of Medical Services in Petroleum Development of Oman

The Head of Medical Services in LNG Oman

ALL PRIVATE PHARMACIES & DRUG STORES

After Compliments,

Please find attached our Circular No 121 dated 29.8.2024 Regarding SFDA Recall of Nasopharyngeal Airway, Double Swivel Connector, 15mm Y Piece, Portex™ Orator Speaking Valve, Thermovent™ 1200 from (mfr: Smiths Medical).

Copy to:

- Director, Office of H.E. The Undersecretary for Health Affairs
- Director of Medical Device Control, DSC
- Director of Pharmacovigilance & Drug Information Dept, DSC
- Director of Drug Control Department, DSC
- Director of Pharmaceutical Licensing Department, DSC
- Director of Central Quality Control Lab., DSC
- Supdt. of Central Drug Information



DSC
مركز سلامة الدواء
Drug Safety Center



ص.ب: ٣٩٣ مسقط - الرمز البريدي: ١٠٠ - هاتف: ٢٢٣٥٧١١١ - فاكس: ٢٢٣٥٨٤٨٩

P.O. Box: 393 Muscat - Postal Code: 100 - Tel: 22357111 - Fax: 22358489

✉ @DSCPHO Email: dscpho@moh.gov.om

URGENT: FIELD SAFETY NOTICE

Nasopharyngeal Airway, Double Swivel Connector, 15mm Y Piece, Portex™ Orator Speaking Valve, Thermovent™ 1200

07 August 2024

Dear Valued Customer,

Smiths Medical is issuing this letter to notify you of a potential issue with the packaging sterile seals on specific Nasopharyngeal Airway, Double Swivel Connector, 15mm Y Piece, Portex™ Orator Speaking Valve, Thermovent™ 1200 products packaged between 20 January 2021 and 27 August 2021. This letter details the issue and the required steps for you to perform.

Issue:

Smiths Medical has identified a timeframe where there is uncertainty in the seal integrity of the sterile packaging of sterilized Nasopharyngeal Airway, Double Swivel Connector, 15mm Y Piece, Portex™ Orator Speaking Valve and Thermovent™ 1200 products.

Potential Risk

The potential risk of the uncertainty in the seal integrity of the sterile packaging is that a product labeled as sterile, may not be sterile, which could potentially lead to infection.

To date, Smiths Medical has received zero (0) complaints or adverse events associated with this issue.

Affected Product

The affected product SKUs and lots, which were manufactured between 20 January 2021 and 27 August 2021 are listed below.

Table 1: Affected Product(s)

SKU	Description	Lot #
100/210/060	Nasopharyngeal Airway 6.0MM 10/BX	4107295 4115880 4125006 4133226 4133228
100/210/070	Nasopharyngeal Airway 7.0MM 10/BX	4089150 4097684 4110350 4130777 4133227 4147167
100/210/080	Nasopharyngeal Airway 8.0MM 10/BX	4107296 4112944 4127687
100/210/090	Nasopharyngeal Airway 9.00MM 10/BX	4133814
100/255/150	15mm Double Swivel Connector 10/BX	4089148

		4107293
100/550/000	Portex "orator" speaking valve for tracheostomy tube 2/CA	4097683 4131840
100/582/000	Thermovent™ 1200-15MM/22MM 20/BX	4088216 4088217

Smiths Medical Actions:

Smiths Medical is sending this notification to all customers who received product(s) from Smiths Medical listed above. Smiths Medical has initiated a global ship hold on impacted lots to ensure affected product is no longer distributed. Smiths Medical will provide credit to affected customers.

Customer Required Actions:

- 1) Check all inventory locations within your institution for the affected catalog numbers and lot numbers listed in the notification and discontinue use. Discard all affected products following your institution's process for discarding. If discarding is not immediately possible at your facility, then the product should be quarantined until disposal.
- 2) Share this notification with all potential users of the device to ensure they are aware of this notification and proposed mitigations. If the devices are used at another location, please ensure this communication is delivered there.
- 3) Complete and return the attached Customer Response Form to EMEA-FSN@icumed.com within 10 days of receipt to acknowledge your understanding of this notification.
- 4) **DISTRIBUTORS:** If you have distributed potentially affected products to your customers, please immediately forward this notice to them and request that they complete the response form and return it to **YOU**. Then the **DISTRIBUTOR** must complete a SINGLE form with the required details and return to EMEA-FSN@icumed.com

For further inquiries, please contact the applicable team using the following information:

Smiths Medical Contact	Contact Information	Areas of Support
Global Complaint Management	globalcomplaints@icumed.com	To report adverse events or product complaints
Field Safety Notice	EMEA-FSN@icumed.com or contact your sales representative	Questions about this Field Safety Notice

Your country regulatory agency has been notified of this action

Smiths Medical is committed to patient safety and is focused on providing exceptional product reliability and the highest level of customer satisfaction. Thank you for your prompt support on this important matter. We appreciate your cooperation.

Sincerely,



Andy Mathein
Vice President of Quality

URGENT: FIELD SAFETY NOTICE – RESPONSE FORM

Nasopharyngeal Airway, Double Swivel Connector, 15mm Y Piece, Portex™ Orator Speaking Valve, Thermovent™ 1200

07 August 2024

Check your inventory and complete the information below, even if you do not have the affected product. *Failure to complete all sections of this page may result in improper, delayed or denied credit.*

Please return the completed form to EMEA-FSN@icumed.com. If you have questions about this form please contact EMEA-FSN@icumed.com or your local sales representative.

Name of Hospital / Facility	
Hospital / Facility Address	
Telephone Number	
Name and Title of Person Completing this Form	
Signature of Person Completing this Form	
Date	
If Purchased through a distributor, please list distributor name/location here for traceability purposes	

Please select one:

- ☐ I have **NO** affected products (complete and return this form to the e-mail address above)
- ☐ **YES**, I have affected products, I have notified users in my facility and I have followed the instructions provided to me and destroyed all affected items (see table below)

If you have affected product on hand, please complete table below:

TABLE 1

Lot Number	Quantity in inventory	Quantity Destroyed	Date of Destruction	PO, debit memo or invoice

If you have distributed the product further, please complete table below with collated information received from your customers and respond to ICU Medical with the overall information.

TABLE 2

Lot Number	Quantity destroyed locally by customer	Date of Destruction

Adverse events and complaints associated with the use of this product should be reported and emailed to Smiths Medical's Global Complaint Management Department at globalcomplaints@icumed.com.

SA-01-08-24-539

Details

Reference Number

SA-01-08-24-539

Manufacturer Company

Smiths Medical ASD, Inc.

Medical Device Name

Nasopharyngeal Airway, Double Swivel Connector, 15mm Y Piece, Portex™ Orator Speaking Valve, Thermovent™ 1200

Affected Devices

Refer to Table 1 in the attachment for affected lots (manufactured between 20 January 2021 and 27 August 2021).

Authorized Representative

AL MADAR MEDICAL SERVICES COMPANY

Reason of Problem

There is uncertainty in the seal integrity of the sterile packaging of sterilized above-mentioned products.

Action

- Discontinue use and discard all affected products following your institution's process for discarding. If discarding is not immediately possible at your facility, then the product should be quarantined until disposal.
- Credit will be provided by Smiths Medical to the affected customers.

Attach Files



Back