

Date: 28-April-2023

Urgent Field Safety Notice Mölnlycke® Barrier TUR Set

For Attention of Theatre Manager

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

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

Name: Local Customer Care contact will be added for each specific market



Date: 28-April-2023

1.

1.

<u>Urgent Field Safety Notice (FSN)</u> <u>Mölnlycke® Barrier TUR Set</u>

1. Information on Affected Devices 1. Device Type(s) Product codes: 888224-22 TUR Set Surgical drape, general/plastic surgical procedure kit, non-medicated, single-use Sterile 1. 2. Commercial name(s) BARRIER TUR Set 1. 3. Primary clinical purpose of device(s) The surgical drape sets consists of both surgical drapes and supplementary products intended be used to create a sterile field for a surgical procedure.

Description of the product problem*

See Appendix I Product Table

See Appendix I Product Table

Molnlycke have identified a composition error in the TUR Set. The incorrect drape has been included within the set. Drape $\underline{965520-22}$ has been included instead of drape $\underline{965522-22}$.

The products have different features and materials connected to the fluid pouch and therefore the dose applied to reach sterility is not high enough.

Reason for Field Safety Corrective Action (FSCA)

As a precaution Mölnlycke has decided to perform a Recall.

4. Device Model/Catalogue/part number(s)

5. Affected serial or lot number range

Hazard giving rise to the FSCA*

If there is a compromise in sterility of drapes which are used during a surgical procedure, there is a potential risk of a local infection.



Date: 28-April-2023

3. Type of Action to mitigate the risk

3. 1. Action To Be Taken by the User

- □ Quarantine Device
- ⊠ Return Device

We need your help in ensuring that <u>all affected products</u> are located and that below actions are performed.

Please follow below instructions:

- 1. **Identify and isolate** the unused Mölnlycke® BARRIER TUR Set at your facility, please see Appendix I for affected product information.
- Fill out the Customer Reply Form or Distributor Reply Form with quantity of identified affected products. Please sign and email/fax the Customer Reply Form or Distributor Reply Form per its instructions within 10 business days.
- Even if you no longer have any concerned Mölnlycke® BARRIER TUR Set, fill
 out the Customer Reply Form or Distributor Reply Form and return it back
 within 10 business days. Mölnlycke needs to be sure all customers are aware of
 the situation.
- Mölnlycke will contact you and arrange for collection of the product(s) from your facility, as soon as you return the Customer Reply Form or Distributor Reply Form. Mölnlycke will issue a credit for the goods returned.
- 5. If you have forwarded any affected products to other healthcare institutions, please send them a copy of this **Field Safety Notice**. Make sure they act accordingly.
- 6. If you are a distributor, please inform your customers by sending them a copy of this **Field Safety Notice**. Make sure they act accordingly and return the **Distributor Reply Form** with information collected from your end users.

We apologize for any inconvenience this will cause you, and rest assured it is our utmost intent to make this process as easy for you as possible.

In addition, Mölnlycke appreciates your help in collecting data on product complaints and/or incidents related to the concerned product. Please follow the reporting procedures established by your facility.

3.	Is customer Reply Required? (If yes, form attached	Yes (Within	10
	specifying deadline for return)	business days)	
l			



Date: 28-April-2023

	4. General Information		
4.	FSN Type	New	
4.	Further advice or information already expected in follow-up FSN?	No	
4.	Manufacturer information For contact details of local representative	contative refer to make 4 of this ECNI	
(For contact details of local representative refer to page 1 of a. Company Name Mölnlycke Hea		Mölnlycke Health Care AB	
	b. Address	Box 130 80, SE-402 52 Gothenburg, Sweden	
	c. Website address	www.molnlycke.com	
4.	The Competent (Regulatory) Authority of your country has been informed about this communication to customers.		
4.	List of attachments/appendices:	Appendix I Product table Customer Reply Form	
4.	Name/Signature		

Transmission of this Field Safety Notice

This notice needs to be passed on 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.



FSCA Ref: 2023-04(01)

FSN Ref: 2023-04(01)
Date: 28-April-2023
Appendix I

Product table

Material	Material Description	Batch
888224-22	TUR Set	23136207



FSN Ref: 2023-04(01) Date: 26 Apr 2023 FSCA Ref: 2023-04(01)

Customer Reply Form

1. Field Safety Notice (FSN) in	formation			
FSN Reference number		2023-04(01)		
FSN Date		26 Apı		
Product/ Device name			ppendix I Product table	
Troduct Borios Harris		0007	pportaix i i roddot tablo	
Product Code(s)		See A	ppendix I Product table	
Batch/Serial Number (s)		See A	ppendix I Product table	
2. Customer Details				
Account Number				
Healthcare Organisation Name*				
Organisation Address*				
Department/Unit				
Shipping address if different to ab	ove			
Contact Name*				
Title or Function				
Telephone number*				
Email*				
3. Customer action undertake	n on behalf o	f Health	ncare Organisation	
☐ I confirm receipt of the Field Safety Notice and that I read and understood its content. I do not have any affected devices.				
☐ I confirm receipt of the Field Safety Notice and that				
I read and understood its	Qty:		Lot/Serial Number:	\neg
content.	N/A		Comments	4
I have quarantined affected devices ready for return -			Comments	
enter number of devices ready for return				
Print Name*				
Signature*				
Date*				
4. Return acknowledgement to	sender			
Email		vigilan	ce@molnlycke.com	
Customer Helpline		+46 20-79 82 64		
Postal Address		Mölnlycke Health Care,		
		Box 130 80, SE-402 52		
		Gothenburg, Sweden		



Date: 26 Apr 2023

Fax	+46 31 722 34 00
Deadline for returning the customer reply form*	Within 10 days

Mandatory fields are marked with *

It is important that your organisation takes the actions detailed in the FSN and confirms that you have received the FSN.

Your organisation's reply is the evidence we need to monitor the progress of the corrective actions.