

Rev 1: September 2018 FSN Ref: CAR196

FSCA Ref: CAR196

Date: 12/06/2025

Urgent Field Safety Notice (RECALL)

Guedel Airways

For Attention of*: MDSO's, All clinical staff, Managers and users of the above products, including those who may use them remotely.

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: <u>giedriusb@intersurgical.lt</u> Tel. +370 387 66611 Fax: +370 387 66622

or

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



Urgent Field Safety Notice (FSN)

Guedel Airways

Risk addressed by FSN

	1. Information on Affected Devices*			
1.	1. Device Type(s)*			
	Guedel Airway			
1.	2. Commercial name(s)			
	One-piece Guedel airway, size 2, ISO 8.0, green One-piece Guedel airway, size 3, ISO 9.0, yellow One-piece Guedel airway, green, ISO 8.0, size 2 (grouped in 10s) One-piece Guedel airway, yellow, ISO 9.0, size 3 (grouped in 10s)			
1.	3. Unique Device Identifier(s) (UDI-DI)			
	5030267050659 5030267050680 5030267091997 5030267091966			
	 Primary clinical purpose of device(s)* 			
	To establish and maintain a patent airway.			
1.	5. Device Model/Catalogue/part number(s)*			
	REF: 1112080 REF: 1113090 REF: 8112080 REF: 8113090			
1.	6. Software version			
	N/A			
1.	7. Affected serial or lot number range			
	REF: 1112080 32407072 32408311 32409113 32409836 32410538 32411087 32413963 32414447 32415284 32415941 32416250 32420438 32420919 32421649 32422162 32422693 32423127			
	32418359 32419010 32419599 32420657 32421079 32422296 32423332 3242384 32424213			



	REF: 8112 32407640	080 32408994	32412180	32414671	32418784	32421509	
	REF: 8113 32406965		32412074	32417555	32418164	32421904	32424085
1.	8. Ass N/A.	ociated devid	ces				

	2. Reason for Field Safety Corrective Action (FSCA)*				
2.	1. Description of the product problem*				
	During manufacture small plastic burrs have been identified inside the Guedel Airways, as shown below.				
2.	2. Hazard giving rise to the FSCA*				
	The device is potentially contaminated with small plastic burrs inside the Guedel or packaging from the manufacturing process. If the burr becomes detached and is inhaled, it could result in potential complications such as airway obstruction, tissue irritation, inflammation and infection.				
2.	3. Probability of problem arising				
	High in the affected Lot number range.				
2.	4. Predicted risk to patient/users The risks associated with the identified fault have been reviewed, where the probability of harm is low, but due to the higher rate of possible occurrence we feel it is essential to address the issue promptly to further reduce the risk of any potential patient harm.				
2.	5. Further information to help characterise the problem				
	N/A				



2.	6. Background on Issue			
	A non-conformance was raised for plastic burrs found inside the Guedel Airway during manufacture.			
	The fault was caused by damaged production equipment contacting the inside of the Guedel Airway,			
	which has since been corrected with no further problems identified. Further evaluation of products			
	and manufacturing records has ena	abled us to confirm the rang	ge of products and lot numbers under	
	this FSCA. No reports of this problem have been reported from the market to-date.			
2.	7. Other information relevant t	o FSCA		
	N/A			
	3. Type of Action to mitigate	the risk*		
3.	1. Action To Be Taken by the U			
	-			
	🛛 Identify Device 🛛 🖾 Quara	ntine Device 🛛 🛛 Return	Device Destroy Device	
	,		y	
	On-site device modification/i	aspection		
		lopoolion		
		recommendations		
	□ Follow patient management	recommendations		
		- f f - f f		
	□ Take note of amendment/rei	nforcement of Instructions	-or Use (IFU)	
	⊠ Other □ None			
			of the Guedel Airway devices listed	
		or their awareness of the po	otential problem and to carry out the	
	following actions.			
	1. Identify and immediately quarantine any potentially affected products from the affected code and			
	lot numbers listed above.			
	2. Please complete the Reply Form below to confirm the products you have identified, to arrange			
	collection of the devices and a credit.			
	3. If you have no affected devices i			
			<u>Dintersurgical.lt</u> , to confirm receipt of	
	this notice and that the necessary actions have been taken.			
	Please note: This is a Recall.			
	Please continue to report to Intersurgical any adverse events involving this product.			
3.		nediately on receipt of this	0 0	
	action be completed? affe	cted stock listed in this FS	N is remaining.	
3.	3. Particular considerations for: N	/Α		
	Is follow-up of patients or review	v of patients' previous resu	Its recommended?	
	Not applicable.			
3.	4. Is customer Reply Required? *		Yes	
	(If yes, form attached specifying de	adline for return)		



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3.	5.	Action Being Taken by the Manufacturer			
		⊠ Product Removal □ Software upgrade □ Other	 □ On-site device modification □ IFU or labelling change □ None 	/inspection	
3	6.	By when should the action be completed?	Two months from receipt o	of the FSN	
3.	7.	Is the FSN required to be communicated to the patient /lay user?		No	
3	8.		provided additional information si sional user information letter/shee	uitable for the patient/lay user in a et?	

	4. General Information*				
4.	1. FSN Type*	New – Recall Notice			
4.	2. For updated FSN, reference	N/A			
	number and date of previous FSN				
4.	3. For Updated FSN, key new information as follows:				
	N/A				
4.	4. Further advice or information	No			
	already expected in follow-up FSN?*				
840	5. If follow-up FSN expected, what is t	he further advice expected to relate to:			
4 N/A					
	6. Anticipated timescale for follow-up	N/A			
4	FSN				
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) Auth communication to customers. *	nority of your country has been informed about this			
4.	9. List of attachments/appendices:	Customer Reply Form			
4.	10. Name/Signature	Ivan Seniut, Group Quality and Regulatory Affairs Director, Intersurgical			



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.

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