

 LeMaitre Vascular GmbH Otto-Volger-Str. 5a/b 65843 Sulzbach/Ts. Germany
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www.lemaitre.com

Field Safety Notice - Ref : CAPA 2025-007 TufTex® Over-the-Wire Embolectomy Catheter Pruitt® Occlusion Catheter Pruitt® Irrigation Occlusion Catheter

Dear Sir, Madam,

An urgent Field Safety Notice (FSN) related to TufTex[®] Over-the-Wire Embolectomy Catheter, Pruitt[®] Occlusion Catheter and Pruitt[®] Irrigation Occlusion Catheter is attached to this cover letter. Please read it carefully, fill out the response form and email it back to <u>recall-UK@lemaitre.com</u>.

We sincerely apologize for the inconvenience and thank you in advance for your cooperation.

Sincerely yours,

LeMaitre Vascular Ltd Helen GOULDING UK Responsible Person



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Field Safety Notice <u>TufTex® Over-the-Wire Embolectomy Catheter</u> <u>Pruitt® Occlusion Catheter</u> <u>Pruitt® Irrigation Occlusion Catheter</u>

For Attention of: Risk Management

Contact details of local representative / Authorized Representative: Hélène Plas (PRRC) LeMaitre Vascular GmbH Otto-Volger-Strasse 5a/b 65843 Sulzbach/Taunus Germany regulatory-emea@lemaitre.com +33 (0)6 75 22 32 16



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Field Safety Notice (FSN) <u>TufTex® Over-the-Wire Embolectomy Catheter</u> <u>Pruitt® Occlusion Catheter</u> <u>Pruitt® Irrigation Occlusion Catheter</u>

1. Information on Affected Devices

1.1. Device Type(s):	Embolectomy Catheter, Occlusion Catheter.		
1.2. Commercial name(s):	TufTex® Over-the-Wire Embolectomy Catheter, Pruitt® Occlusion		
	Catheter, Pruitt® Irrigation Occlusion Catheter		
1.3. Unique Device Identifier(s)	1651-34: 00840663100651		
(UDI-DI):	1651-38: 00840663100668		
	1651-44: 00840663100675		
	1651-48: 00840663100682		
	1651-64: 00840663100712		
	1651-68: 00840663100729		
	1651-78: 00840663100736		
	1651-84: 00840663100743		
	1651-88: 00840663100750		
	2103-36: 00840663101559		
	2103-46: 00840663101566		
	2103-56: 00840663101573		
	2102-09: 00840663101535		
1.4. Primary clinical purpose of	TufTex® Over-the-Wire Embolectomy Catheter: indicated for use in		
device(s):	the removal of emboli and thrombi during embolectomy and/or		
	thrombectomy. It can also be used for catheter placement over a		
	guidewire, vessel occlusion, fluid infusion and/or aspiration.		
	Pruitt® Occlusion Catheter: indicated for the occlusion of vessels		
	both arterial and venous for the control of bleeding.		
	Pruitt® Irrigation Occlusion Catheter: indicated to temporarily oc-		
	clude vessels for the control of bleeding. To access the vessel lu-		
	men distal to the point of occlusion.		
1.5. Device Model/	TufTex® Over-the-Wire Embolectomy Catheter: 1651-34, 1651-38,		
Catalogue / part number(s)	: 1651-44, 1651-48, 1651-64, 1651-68, 1651-78, 1651-84, 1651-88		
	Pruitt® Occlusion Catheter: 2103-36, 2103-46, 2103-56,		
	Pruitt® Irrigation Occlusion Catheter: 2102-09		
1.6. Affected serial or lot number	r See attached Impacted Lots List document.		
range:	The potentially impacted lot numbers were determined by reviewing		
	the maximum shelf life of each product line.		
	TufTex® Over-the-Wire Embolectomy Catheter (6 years shelf life) -		
	distributed from April 2019 to April 2025		
	Pruitt® Occlusion Catheter (7 years shelf life) - distributed from July		
	2018 to April 2025		
	Pruitt® Irrigation Occlusion Catheter (5 years shelf life) – distributed		
	from September 2020 to April 2025		

2. Reason for Field Safety Corrective Action (FSCA)

2.1. Description of the product prob-	Inadequate Tray Seals were found on samples from these
lem:	three product lines.





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2.2. Hazard giving rise to the FSCA:2.3. Probability of problem arising:	The potential hazard is infection if the sterile barrier of the
2.3. Probability of problem arising:	packaged device is broken.
	No complaints have been received about this issue. Following inspection and testing of finished product samples, the results suggest that only approx. 1% of the population may have a compromised package.
2.4. Predicted risk to patient / users:	The potential hazard is infection to the patient if the sterile bar- rier of the packaged device is broken. Only patients undergoing surgery with the use of the TufTex® Over-the-Wire Embolec- tomy Catheters, Pruitt® Occlusion Catheters, or Pruitt® Irriga- tion Occlusion Catheters are at risk.
2.5. Further information to help char- acterize the problem:	None
2.6. Background on Issue:	The inadequate tray seal issue was observed during an unre- lated test to TufTex® OTW Catheters. It was then further re- viewed and confirmed by Quality and Manufacturing Engineer- ing where some parts (roughly 10% of any given lot) could ex- hibit voids on the seal area in the location where the tubing connects to the boat tray. The voids were observed visually against backlight and confirmed upon opening of Tyvek lid. Seal integrity testing (dye penetration) of other samples with visually detected voids subsequently confirmed failures in some of the samples, indicating compromised packaging (1%), which can allow microorganisms to enter and contaminate the device, leading to potential infection transmitted by the compro- mised device.
3. Type of Action to mitigate the risk	
3.1. Action To Be Taken by the	☑ Identify Device
User:	⊠ Return Device □ Destroy Device
	□ On-site device modification/inspection
	□ Follow patient management recommendations
	□ Take note of amendment/reinforcement of Instructions For Use (IFU)
	 □ Take note of amendment/reinforcement of Instructions For Use (IFU) □ Other □ None
	 Take note of amendment/reinforcement of Instructions For Use (IFU) Other None Quarantine the product. Complete the form at the end of the FSN and return the form to LeMaitre Vascular GmbH.
3.2. By when should the action be completed (by the user)?	 Take note of amendment/reinforcement of Instructions For Use (IFU) Other None Quarantine the product. Complete the form at the end of the FSN and return the form to LeMaitre Vascular GmbH. As soon as possible.
	 □ Take note of amendment/reinforcement of Instructions For Use (IFU) □ Other □ None Quarantine the product. Complete the form at the end of the FSN and return the form to LeMaitre Vascular GmbH. As soon as possible. Is follow-up of patients or review of patients' previous results recommended? □ Yes ⊠ No
completed (by the user)?	 □ Take note of amendment/reinforcement of Instructions For Use (IFU) □ Other □ None Quarantine the product. Complete the form at the end of the FSN and return the form to LeMaitre Vascular GmbH. As soon as possible. Is follow-up of patients or review of patients' previous results recommended? □ Yes ☑ No No follow up action is required.
completed (by the user)? 3.3. Particular considerations for: 3.4. Is customer Reply Required?	 □ Take note of amendment/reinforcement of Instructions For Use (IFU) □ Other □ None Quarantine the product. Complete the form at the end of the FSN and return the form to LeMaitre Vascular GmbH. As soon as possible. Is follow-up of patients or review of patients' previous results recommended? □ Yes ⊠ No No follow up action is required. ⊠ Yes □ No
3.3. Particular considerations for:	 □ Take note of amendment/reinforcement of Instructions For Use (IFU) □ Other □ None Quarantine the product. Complete the form at the end of the FSN and return the form to LeMaitre Vascular GmbH. As soon as possible. Is follow-up of patients or review of patients' previous results recommended? □ Yes ☑ No No follow up action is required.

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FSN / FSCA Ref: CAPA 2025-007

By when should the action be completed (by the manufac- turer)?	30 October 2025			
eneral Information				
FSN Type:	New			
Further advice or information al- ready expected in follow-up FSN?	□ Yes	□ No	⊠ Not planned yet	
4.3. Manufacturer information:	(For contact detail this FSN)	ls of local rep	presentative refer to page 1 of	
	Company Name:	LeMaitre Vascular, Inc.		
	Address:	63 Second	Ave. Burlington,	
		MA 01803		
		USA		
	Website address:	www.lemait	re.com	
The Competent (Regulatory) Auth tion to customers.	ority of your country	/ has been in	formed about this communica	
	completed (by the manufac- turer)? eneral Information FSN Type: Further advice or information al- ready expected in follow-up FSN? Manufacturer information: The Competent (Regulatory) Auth	completed (by the manufacturer)? eneral Information FSN Type: New Further advice or information already expected in follow-up □ Yes FSN? (For contact detail this FSN) Manufacturer information: (For contact detail this FSN) Company Name: Address: Website address: Website address:	completed (by the manufacturer)? eneral Information FSN Type: New Further advice or information already expected in follow-up Yes No FSN? Manufacturer information: (For contact details of local reprthis FSN) Manufacturer information: Company Name: LeMaitre Va Address: 63 Second MA 01803 USA Website address: www.lemait The Competent (Regulatory) Authority of your country has been in	

4.5. Name / Signature	Helen GOULDING,	
no: Hame / eignatare		
	UK Responsible Person	

5. Transmission of this Field Safety Notice

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

Please transfer this notice to other organizations 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.





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Customer Reply Form

Please complete this reply form and e-mail it to us at <u>recall-UK@lemaitre.com</u>. **The form must be returned even if you have zero devices in inventory.**

Account #	Customer Name	Address	
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If you are not the customer listed here, please list your facility information below.

Contact Name			
(First and Last Name)			
Contact Email			
Contact Phone			
Signature and Date			
Do you have any recalled devices at your facility? If Yes, please complete the table below.	□Yes	🗆 No	

• If you have checked your inventory and have no recalled devices, you may simply email recall-UK@lemaitre.com to indicate that "I have checked our inventory and we have none of the recalled devices."

REF #	LOT #	QUANTITY ON HAND

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ADDRESS TO WHICH REPLACEMENT DEVICES SHOULD BE SENT:

Distributors:

 \Box I have checked my stock and have quarantined inventory consisting of _____ units.

 \Box I identified and notified all of my customers that are affected by this recall.

 \Box If the product was distributed outside the US, I have notified that country's medical device regulatory agency about this recall.

 \Box I did not notify the regulatory agency. The rationale is listed below.

Rationale:

Name/Title

Telephone

Email address

If you have transferred devices to another facility, please send them a copy of this recall letter. If possible: list the facility information, including contact information. Also, please add a note if you received the devices from another facility.

Thank you for your cooperation!

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