

7th November 2016

URGENT FIELD SAFETY NOTICE

Commercial Name of Affected Product:		ARROW® OnControl® Bone Lesion Biopsy System Tray ARROW® OnControl® Aspiration System Tray ARROW® OnControl® Bone Lesion Biopsy System Tray ARROW® OnControl® Bone Lesion Biopsy System Trays ARROW® OnControl® Bone Marrow Biopsy System Comprehensive Tray ARROW® OnControl® Bone Marrow Biopsy System Tray ARROW® OnControl® Ported Aspiration System Tray ARROW® OnControl® System Sterile Procedure Tray OnControl® Biopsy System Ported Needle Tray			
Type of action:		Recall			
Teleflex Reference:		EIF-000084			
Product Code	Lot/Batch	Product Code	Lot/Batch	Product Code	Lot/Batch
9403-EU-006	Refer to Appendix 2	9461-VC-006	Refer to Appendix 2	9464-VC-006	Refer to Appendix 2
9403-VC-006		9462-EU-001		9465-VC-001	
9408-EU-006		9462-VC-001		9465-VC-006	
9408-VC-006		9462-VC-006		9466-EU-001	
9411-EU-006		9463-EU-001		9466-VC-001	
9411-VC-006		9463-VC-001		9466-VC-006	
9451-VC-006		9463-VC-006		9470-VC-006	
9458-VC-006		9464-EU-001		9471-VC-006	
9461-VC-001		9464-VC-001		9472-VC-006	

Dear Customer,

Details of affected devices

Teleflex has initiated a voluntary Field Safety Corrective Action for the above listed products.

Description of the problem

Teleflex is recalling this product due to a potential incomplete seal on the outer sterile package. Because of the compromised packaging, the sterility of the inside drape, which is used in preparation for bone marrow aspiration with the OnControl system, cannot be guaranteed. If sterility of the drape is compromised, there is a potential for infection to occur.

FIELD SAFETY CORRECTIVE ACTION INSTRUCTIONS

ADVICE ON ACTION TO BE TAKEN BY MEDICAL STAFF

1. We request that you check your inventory for product within the scope of this field action. Users should cease use and distribution of stock of affected product and quarantine immediately.
2. If you do not have stock in scope of this field action as referred to in above table, then mark the according checkbox on the Acknowledgement form (Appendix 1) and return the form to the fax number or e-Mail address mentioned there.



3. If you have stock from the affected product, mark the according checkbox on the Acknowledgement form (Appendix 1). Contact customer service by calling the phone number mentioned below, who will issue you with a return number. Write this return number into the respective field in the Acknowledgement form.
4. Complete 'Appendix 1' for all products in your possession and under control. Return this form immediately to the fax number below or provide a completed copy to your local Sales Representative.
5. Teleflex (or your local dealer) will issue a credit note upon receipt of the returned affected product.

INSTRUCTION FOR DISTRIBUTORS OF AFFECTED PRODUCT

1. Immediately discontinue distribution and quarantine any products with the catalog and lot number listed above.
2. Using the provided customer letter and Recall Acknowledgement Form templates, communicate this recall to any of your customers who have received product included within the scope of the recall.
3. Have the customers return any affected product to you, together with a completed Recall Acknowledgement Form, for consolidation and return to Teleflex. In the event that an alternative approach is needed, contact Teleflex Customer Service for more information.
4. To return product, complete the enclosed Recall Acknowledgement Form and email or fax it to the contact listed on this notice. This will allow us to document the amount of product you have on hand for return. A customer service representative will contact you with a Return Goods Authorization (RGA) Number and will provide instructions for the return of product to Teleflex Medical.
5. If you and your customers have no affected stock, please complete the enclosed Recall Acknowledgement Form and email or fax it to the contact listed on this notice. This will allow us to document your receipt of this letter.
6. Once you have completed returning all of the recalled products from your own inventory, and collecting and consolidating all of the recalled products from your customers, please check the box on the enclosed Recall Acknowledgement Form that indicates that you have completed the recall and email or fax it to the contact listed on this notice. This will allow us to document completion of the recall.
7. Please be aware that all European Economic Area/Switzerland (EEA/CH) and Turkey Member State Competent Authorities in which Teleflex distribute directly will be notified by Teleflex.
8. If you are a distributor and/or have a reporting responsibility within or outside the EEA/CH/TK area, please notify your local Competent Authority of this action. Please forward the notification and all communication with your local competent authority to Teleflex.

Teleflex

Teleflex informs all customers, employees of Teleflex and distributors on this Field Action.

Transmission of this Field Safety Notice

This notice should be passed on all persons who need to be aware within your organization or to any organization where the potentially affected devices have been transferred. Please consider end users, clinicians, risk managers, supply chain/distribution centres etc. in the circulation of this notice. Maintain awareness of this notice until all required actions have been completed in your organisation

Contact reference person

Should you require any further information or support concerning this issue, please contact:



Customer Service

Contact: Shane Kenny
FAX: +353 (0)1 4370773

Telephone: +353 (0)906460869
E-mail: Recalls.intl@teleflex.com

Teleflex is committed to providing high quality, safe and effective products. We sincerely apologize for any inconvenience this action may cause your operations. If you have any other questions, feel free to contact your local sales representative or Customer Service.

For and on behalf of Teleflex,

Karen Boylan

Karen Boylan - VP, Global RA/QA



Appendix 1

Customer No: _____

FIELD SAFETY CORRECTIVE ACTION

Teleflex Ref. EIF-000084

Acknowledgement Form

URGENT ATTENTION REQUIRED

Return completed form immediately to:

FAX: +353 (0)1 4370773

E-mail: Recalls.intl@teleflex.com

Please check applicable box:

<input type="checkbox"/> We confirm receipt of this FSN and completed the required actions contained therein. We confirm that our inventory does NOT include products affected by this Field Action.	<input type="checkbox"/> We confirm receipt of this FSN and completed the required actions contained therein. We confirm our inventory DOES include products affected by this Field Action. The use and further distribution of the affected products has been stopped. All products are on hold and the quantity stated below will be returned.
<div style="border: 2px solid red; padding: 5px; display: inline-block;"> Return Authorisation No _____ </div>	

Please CLEARLY print the below return information

Name of Affected Products	ARROW® OnControl® Bone Lesion Biopsy System Tray ARROW® OnControl® Aspiration System Tray ARROW® OnControl® Bone Lesion Biopsy System Tray ARROW® OnControl® Bone Lesion Biopsy System Trays ARROW® OnControl® Bone Marrow Biopsy System Comprehensive Tray ARROW® OnControl® Bone Marrow Biopsy System Tray ARROW® OnControl® Ported Aspiration System Tray ARROW® OnControl® System Sterile Procedure Tray OnControl® Biopsy System Ported Needle Tray	
Product Number	Lot Number	Quantity (Returning)

Return Instructions:

- Please label product returns as "Field Action Returns".
 - Include a copy of this form (including RAN Number) with product returns.
- Returns excluding ALL necessary documentation CANNOT be processed.

Institution Name - (Hospital, Health Care Organisation, etc.)	
Institution Address:	Email Address:
	Phone Number:
Form completed by:	
Print Name:	Institution Stamp:
Signature:	
Date:	

Product Code	Lot/Batch	Product Code	Lot/Batch	Product Code	Lot/Batch	Product Code	Lot/Batch	Product Code	Lot/Batch
9403-EU-006	010742	9408-VC-006	013257	9451-VC-006	012690	9458-VC-006	014132	9464-EU-001	012134S
	011231		013258		013071		014193		013008S
	011942		013477		013247		014194		013067S
	013606		013478		013470		014195		013475S
9403-VC-006	010848		013688		013692		014196		013510S
	011230		013894		013735		014313		013601S
	011587		013895		013898		014314		013691S
	012101		013896		014125		014315		014320S
	012348		013897		014138		014402	014563S	
	012697		014111		014232		014404	014809S	
	013245		014112		014312		014405	9464-VC-001	010744S
	013888		014113		014565		014406		010956S
014127	014114		010561		014607		011718S		
014208	014115		010562		014608		012135S		
014407	014134		010723		014609		012695S		
9408-EU-006	010740		014135		010781		012579S		9462-EU-001
	010948	014136	010782	013469S	010956				
	011049	014199	010817	011227	011051				
	011377	014200	010872	011492	011491				
	011588	014201	010893	011938	011590				
	012117	014202	010952	012694	011719				
	012961	014306	010953	013065	011939				
	013054	014307	010964	013686	012136				
	013476	014308	010969	014615	012350				
	014124	014420	010977	010894S	012696				
	014137	014422	011039	011715S	012778				
	014167	014423	011040	012458	013252				
	014233	014569	011041	013244S	013902				
	014424	014570	011060	013685S	014025				
	014455	014571	011061	013690S	014120				
	014701	014703	011223	014166S	014231				
9408-VC-006	010438	014805	011378	014205S	9462-VC-006	014231			
	010439	010741	011379	014623S		014319			
	010556	010950	011555	014627S		014453			
	010557	011376	011556	010743S		014618			
	010737	012116	011557	010778S		9465-VC-001	010957S		
	010738	012960	011710	010954S			011940S		
	010739	013051	011711	012580S			014321S		
	010946	013687	011712	012891S			9465-VC-006	010957	
	010947	014311	011949	010743		011496			
	011037	014454	011950	010778		011941			
	011046	010440	011951	010954		012137			
	011047	010559	012143	011495		012351			
	011048	010949	012144	011545		013286			
	011215	011052	012145	011582		013901			
	011216	011218	012146	011714		014322			
	011217	011374	012344	012130		9466-EU-001	010897S		
011380	011386	012345	012581	011451S					
011381	011720	012346	013251	013049S					
011383	011948	012460	013860	013846S					
011384	012352	012461	013893	013892S					
011385	012459	012462	014128	014164S					
011675	012578	012575	014316	014207S					
011676	012689	012576	010895S	014430S					
011945	012894	012577	011449S	014624S					
011946	013060	012691	013050S	9466-VC-001	010844S				
011947	013246	012692	014165S		010958S				
012139	013468	012693	014206S		011722S				
012140	013509	012897	014318S		012685S				
012141	013689	012898	014426S	9466-VC-006	010844				
012142	013891	013072	014626S		010958				
012353	014121	013086	010331S		011512				
012354	014139	013248	010955S		011544				
012355	014168	013249	011716S	011589					
012356	014197	013250	012133S	011723					
012463	014203	013471	013066S	012129					
012464	014310	013472	010331	012466					
012465	014425	013473	010955	013887					
012570	014617	013694	011513	014123					
012572	010183	013695	011584	014209					
012573	010560	013696	011717	014323					
012686	010780	013734	011834	9470-VC-006	010859				
012687	010951	013904	012131		011511				
012688	010972	013905	012349	9471-VC-006	013107				
012895	011042	013906	013070		013260				
012896	011050	013907	013474		013889				
013074	011348	014116	013890		014126				
013084	011382	014117	013899	014324					
013085	011583	014118	014230	9472-VC-006	013108				
013122	011709	014119	014317		013261				
013185	011952	014129	014616						
013255	012343	014130	010896S						
013256	012568	014131	011450S						