
Urgent Medical Device Recall Dual Path Guide

July 30, 2018

Attention: Ultrasound Department

This is to inform you of a product recall involving the products labeled as identified below:

676-144 - Needle Guide with covers for use with Siemens BP9-4 transducers

676-150 - Needle Guide for use with Siemens BP9-4 transducers

Lot numbers:

676-144

101000	100=000			
A018067	A025606	A029383	A035568	A040839
A021026	A026609	A032036	A037150	A041444
A021226	A026610	A033139	A037231	A041723
A022447	A028159	A033153	A037585	
A023876	A028645	A035009	A039417	
A025137	A029293	A035010	A040376	

676-150

A035054	A042207				

Dear Valued Customer,

CIVCO has recently discovered a quality issue with dual path needle guide for use with the Siemens BP9-4 transducer. You are being notified, as your facility has been identified as having received product from the affected manufacturing lots. .

Description of the Problem:

The assembly of the needle guides were found to be either missing adhesive or did not have adequate adhesive to hold the assembled cannula in place on the needle guide.



Needle guide assembled



Bond failure

Action to be Taken by the Customer:

CIVCO requests return of all needle guides received from the affected lots identified on the box.

To return the bracket and schedule replacement contact your Account Representative by phone at 800-
445-6741 or 319-248-6757 or via email at <u>order@civco.com</u> .
<u>Transmission of this Recall Notification:</u> This notice is being communicated to all customers who have purchased a kit in which the needle guide is contained from affected lots. CIVCO requests you provide this notification to the appropriate personnel within your facility. If the affected product was distributed outside of your organization, please notify those locations down to the medical facility level.
Thank you for your cooperation and communication on this issue. Please feel free to contact me if you have any questions or concerns regarding the above information.
Sincerely,
James Leong Regulatory Manager CIVCO Medical Solutions
Urgent Medical Device Recall Dual Path Guide

Return By: 17 August 2018

I have been informed of the reguide.		_	ual path needle			
This notice has been read and	understood by the respons	ible party involved.				
1 I have reviewe	I have reviewed my inventory for the affected products and no longer have any in stock.					
2 I have reviewe Number to return the p	ed my inventory for the affectoroduct.	ted products and I need a F	Return Authorization			
List Lot Numb	er List # units	List Lot Number	List # units			
3 Other, please	provide details.					
	_					
Facility Name						
Street Address	City, State	Phone	_			
Print Name / Title	Signature	Date	_			
Please return this certification Attn: Infiniti Bracket at 319-24		email to order@civco.com				
Please contact a CIVCO Cust Customer Consultant by callin 248-6757 between 7:00AM an	g CIVCO Medical Solutions	Multi-Modality Imaging, at	800-445-6741 or 319			
Thank You.						
	vice Recall (OEM / Dist of CIVCO brains		and Biopsy			

Dear Sir or Madam,

Enclosed please find a notification letter for the dual path guide. We have identified that you have distributed kits which include the needle guide from the affected lots listed we have detailed.

This recall is being conducted to the medical facility / user level. We ask that you forward a copy of this notification letter to all customers who have received this product.

Please contact your Account Representative concerning the replacement instructions if you have any questions. You may reach your Representative by calling CIVCO Medical Solutions, Multi-Modality Imaging, at 800-445-6741 or 319-248-6757 between 7:00AM and 5:30PM CST. You may also contact us via email at order@civco.com

We appreciate your assistance with this matter.

Sincerely,

Jim Leong Regulatory Affairs Manager CIVCO Medical Solutions