

Rev 2: February 2020

FSN Ref: VIG-18-2024-FN07 FSCA Ref: VIG-18-2024-FC08

Date: 2023:12:31

Field Safety Notice Amecath Short Term Haemodialysis Catheter Kit

For Attention of*:STG ELISABETH -TWEESTEDEN ZIEKENHU

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

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Field Safety Notice (FSN) Device Commercial Name Risk addressed by FSN

1. Information on Affected Devices*			
1.	1. Device Type(s)*		
	AMECATH Short Term Haemodialysis Catheter is a soft radiopaque biocompatible polyurethane sterile catheter. The catheters are placed centrally or through femoral vein. Intended Use: Sterile single use device indicated for use in attaining short term access for Haemodialysis or aphaeresis. Supplied Sterile		
1.	2. Commercial name(s)*		
	AMECATH Short Term Haemodialysis Catheter		
1.	Unique Device Identifier(s) (UDI-DI)		
	6221139DIA-SHT-2aXT		
1.	4. Primary clinical purpose of device(s)*		
	indicated for use in attaining short term access for Haemodialysis or aphaeresis. Supplied Sterile		
1.	5. Device Model/Catalogue/part number(s)*		
	SDLC-1415-KPC & all SDLC range distributed in 2023 and 2024		
1.	6. Software version		
	NA		
1.	7. Affected serial or lot number range		
	Batch: 510515, Manufacturing date: 03/02/2023 Expiration Date: 31/01/2026		
1.	Associated devices		
	NA		

	2. Reason for Field Safety Corrective Action (FSCA)*				
2.	2. 1. Description of the product problem*				
	Unfortunately we have received a new complaint about the same issue of the suture-wing on an SDLC catheter dislodging from the catheter, resulting in the catheter (partially) existing the blood vessel.				
2.	2. Hazard giving rise to the FSCA*				
Catheter inexplicably removed from patient after insertion. It appears the sutur					
slipped over its hub sometime after insertion					
2.	Probability of problem arising				
This case was encountered with our customer in Netherlands and was handled .					
4. Predicted risk to patient/users					
Evaluated as Critical and it may cause a harm on the patient. Evaluated as critical,					
	mobility of the fixation wing (which is crucial to fix the catheter in place) may lead to				
	catheter slippage out of the vessel, which may cause bleeding and put patient's healt				
	risk.				
2.	5. Further information to help characterise the problem				
	NA				
2.	6. Background on Issue				
	Manufacturer became aware when the distributor notified us by mail Root cause of the				
	problem is missing the inspection step on the assembly step of the rotating wing over the				
hub. The containment action is to create FSNs to be circulated to users who					
defect batch, and they will be instructed to use "Unifix Catheter Tube Fixation					



Rev 2: February 2020

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for catheter fixation instead of rotating wing suturing. In addition, to prevent the incident recurrence, a step will be added in visual inspection process to ensure the proper assembly of the rotating wing over the hub and to ensure that the rotating wing is placed after the stopper. In addition, a process step was edited by changing the mold used to manufacturer these catheters. Then, a recall of defective lots was suggested by the Dutch HA, consequently the manufacturer followed by applying a voluntary recall on the whole SDLC range supplied during the year 2023 and 2024 till date.

2. 7. Other information relevant to FSCA

"Unifix Catheter Tube Fixation Adhesive" is already included in all supplied kits

	3. Type of Action to mitigate the risk*					
3.	1. Action To Be Taken by the User*					
		☐ Identify Device ☐ Quara	ntine Device ☐ Return Device ☐ Destroy Device			
	☐ On-site device modification / inspection					
		☐ Follow patient management recommendations				
		\square Take note of amendment / reinforcement of Instructions For Use (IFU)				
	⊠ Other □ None					
		SDLC range that was supplied Distributor	d during the year 2023 and 2024 till date will be collected by the			
3.	2.	By when should the action be completed?	Within 45 days from the date of circulation of this FSN			
3.	3.	Particular considerations for	or: Choose an item.			
		Is follow-up of patients or review of patients' previous results recommended? Choose an item.				
		Provide further details of patient-level follow-up if required or a justification why none is required.				
3.	(If	4. Is customer Reply Required? * (If yes, form attached specifying deadline for return)				
3.	5 .	Action Being Taken by	the Manufacturer*			
		☑ Product Removal☐ Software upgrade☐ Other	☐ On-site device modification/inspection☐ IFU or labelling change☐ None			
		a recall of defective lots was suggested by the Dutch HA, consequently the manufacturer followed by applying a voluntary recall on the whole SDLC range supplied during the year 2023 and 2024 till date				
3.	6.	By when should the	Within 45 working days. Action is not critical to			



Rev 2: February 2020

FSN Ref: VIG-18-2024-FN07 FSCA Ref: VIG-18-2024-FC08

3.	7.	Is the FSN required to be communicated to the patient No		
		/lay user?		
3.	8.	If yes, has manufacturer provided additional information suitable for the patient/lay		
	user in a patient/lay or non-professional user information letter/sheet?			
		Choose an item. Choose an item.		

4. General Information*				
4.	1. FSN Type*	New		
4.	For updated FSN, reference number and date of previous FSN	Provide reference and date of previous FSN if relevant.		
4.	3. For Updated FSN, key new information as follows:			
	Summarise any key difference in devices affected and/or action to be taken.			
4.	4. Further advice or information already expected in follow-up FSN? *	No		
4.	5. If follow-up FSN expected, what is the further advice expected to relate to:			
Eg patient management, device modifications etc.				
4.	6. Anticipated timescale for follow- up FSN	For provision of updated advice.		
4.	7. Manufacturer information (For contact details of local representative refer to page 1 of this FSN)			
	a. Company Name	Ameco Medical Industries		
	b. Address	Industrial area B4 Plot 119 east, 10th of		
		Ramadan city - Egypt		
	c. Website address	www.amecathgroup.com		
4.	8. The Competent (Regulatory) Authority of your country has been informed about this communication to customers. *			
4.	9. List of attachments/appendices:	NA		
4.	10. 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.*



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