

Rev 1: September 2018
FSN Ref: FSN-2022-010

FSCA Ref: FSN-2022-010

Date: 20 September 2022

Urgent Field Safety Notice

COLUMBIA CAP WITH SHEEP BLOOD – PB5082A, Lot 3532554

For Attention of*: Lab Managers

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

E.mail : mbd.vigilance@thermofisher.com

Telephone: +44(0) 1256 841144

Fax: +44(0) 1256 479525

Urgent Field Safety Notice (FSN)

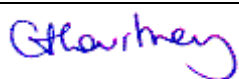
COLUMBIA CAP WITH SHEEP BLOOD – PB5082A, Lot 3532554

1. Information on Affected Devices*	
1.	1. Device Type(s)* Prepared Microbial Culture Media
1.	2. Commercial name(s) Columbia CAP Agar with Sheep Blood
1.	3. Unique Device Identifier(s) (UDI-DI) 5032384128471
1.	4. Primary clinical purpose of device(s)* A selective medium for the isolation of gram-positive bacteria from clinical specimens.
1.	5. Device Model/Catalogue/part number(s)* PB5082A
1.	6. Software version N/A
1.	7. Affected serial or lot number range Lot 3532554
1.	8. Associated devices None

2. Reason for Field Safety Corrective Action (FSCA)*	
2.	1. Description of the product problem* A technical investigation has concluded that Proteus spp grew and swarmed on the plate.
2.	2. Hazard giving rise to the FSCA* Potential not to be able to isolate colonies of target organisms, resulting in delay to treatment.
2.	3. Probability of problem arising High
2.	4. Predicted risk to patient/users Very Low to negligible
2.	5. Further information to help characterise the problem None

2.	6. Background on Issue
	A technical investigation following complaints have concluded that Proteus spp grew and swarmed on the plate. Additional lots have been tested and found to be performing as intended.
2.	7. Other information relevant to FSCA
	Lot number 3532554, Expiry Date - 05 th October 2022

3. Type of Action to mitigate the Risk*		
3.	1. Action To Be Taken by the User*	
	<input checked="" type="checkbox"/> Identify Device <input type="checkbox"/> Quarantine Device <input type="checkbox"/> Return Device <input checked="" type="checkbox"/> Destroy Device <input type="checkbox"/> On-site device modification/inspection <input checked="" type="checkbox"/> Follow patient management recommendations <input type="checkbox"/> Take note of amendment/reinforcement of Instructions For Use (IFU) <input type="checkbox"/> Other <input type="checkbox"/> None	
3.	2. By when should the action be completed?	Immediately
3.	3. Particular considerations for: IVD	
	Is follow-up of patients or review of patients' previous results recommended? Yes Provide further details of patient-level follow-up if required or a justification why none is required	
3.	4. Is customer Reply Required? * (If yes, form attached specifying deadline for return)	Yes
3.	5. Action Being Taken by the Manufacturer	
	<input checked="" type="checkbox"/> Product Removal <input type="checkbox"/> On-site device modification/inspection <input type="checkbox"/> Software upgrade <input type="checkbox"/> IFU or labelling change <input type="checkbox"/> Other <input type="checkbox"/> None	
3	6. By when should the action be completed?	Immediately
3.	7. Is the FSN required to be communicated to the patient /lay user?	No
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?	
	N/A	

4. General Information*		
4.	1. FSN Type*	New
4.	2. For updated FSN, reference number and date of previous FSN	N/A
4.	3. For Updated FSN, key new information as follows:	
	N/A	
4.	4. Further advice or information already expected in follow-up FSN? *	Not planned yet
4	5. If follow-up FSN expected, what is the further advice expected to relate to:	
	N/A	
4	6. Anticipated timescale for follow-up FSN	N/A
4.	7. Manufacturer information (For contact details of local representative refer to page 1 of this FSN)	
	a. Company Name	Oxoid Deutschland GmbH
	b. Address	Am Lippeglacis 4-8 46483, Wesel Germany
	c. Website address	www.thermofisher.com/microbiology
4.	8. The Competent (Regulatory) Authority of your country has been informed about this communication to customers. *	
4.	9. List of attachments/appendices:	Customer Response Form
4.	10. Name	Carissa Courtney Director, Quality EMEA
	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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Customer Reply Form

1. Field Safety Notice (FSN) information			
FSN Reference number*	2022-010		
FSN Date*	20 September 2022		
Product/ Device name*	Columbia CAP Agar with Sheep Blood		
Product Code(s)	PB5082A		
Batch/Serial Number (s)	3532554		
2. Customer Details			
Account Number			
Organisation Name*			
Organisation Address*			
Department/Unit			
Shipping address if different to above			
Contact Name*			
Title or Function			
Telephone number*			
Email*			
3. Customer action undertaken on behalf of Healthcare Organisation			
<input type="checkbox"/>	I confirm receipt of the Field Safety Notice and that I read and understood its content.		
<input type="checkbox"/>	I performed all actions requested by the FSN.		
<input type="checkbox"/>	The information and required actions have been brought to the attention of all relevant users and executed.		
<input type="checkbox"/>	I have returned affected devices - enter number of devices returned and date complete or N/A	Qty:	Lot/Serial Number: Date Returned (DD/MM/YY)
		Comments:	
<input type="checkbox"/>	I have destroyed affected devices – enter number destroyed and date complete.	Qty:	Lot/Serial Number: Date Returned (DD/MM/YY)
		Qty	Credit <input type="checkbox"/> Replacement <input type="checkbox"/>
		Comments:	
<input type="checkbox"/>	No affected devices are available for return/ destruction		
<input type="checkbox"/>	Other Action (Define):		
<input type="checkbox"/>	I do not have any affected devices.		
<input type="checkbox"/>	I have a query please contact me (e.g. need for replacement of the product).		
Print Name*			
Signature*			
Date*			
4. Return acknowledgement to sender			
Email	MBD.vigilance@thermofisher.com		
Telephone Number & Fax	Tel : +44(0) 1256 841144 Fax :+44(0) 1256 479525		
Postal Address			
Deadline for returning the reply form*	18 October 2022		

Mandatory fields are marked with *

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It is important that your organisation takes the actions detailed in the FSN and confirms that you have received the FSN.
Your organisation's reply is the evidence we need to monitor the progress of the corrective actions.

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