Rev 2: February 2020

FSN Ref: FSN2025-01 FSCA Ref: FA-0000102

Date: 10 Jun 2025

Field Safety Notice ACUVUE® OASYS MAX 1-Day MULTIFOCAL

For Attention of: name of the Customer

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

Local country contact details

FSCA Ref: FA-0000102

Field Safety Notice (FSN) ACUVUE® OASYS MAX 1-Day MULTIFOCAL URGENT: MEDICAL DEVICE FIELD SAFETY CORRECTIVE ACTION

1. Information on Affected Devices		
1.	Device Type(s)	
	Soft corrective contact lens, daily-disposable, sterile.	
1.	2. Commercial name(s)	
	ACUVUE® OASYS MAX 1-Day MULTIFOCAL	
1.	3. Unique Device Identifier(s) (UDI-DI)	
	0733905a00011BK	
1.	Primary clinical purpose of device(s)	
	Vision correction	
1.	Device Model/Catalogue/part number(s)	
	888290919116	
1.	6. Affected lot number range	
	J003Q72 1D MAX MULTIFOCAL 8.4 -2.00 LOW 30P RX	

2. Reason for Field Safety Corrective Action (FSCA)

2. 1. Description of the product problem

We are voluntarily initiating a recall of the above referenced lots of ACUVUE® OASYS MAX 1-Day MULTIFOCAL contact lenses due to a defect that may lead to unexpected visual disturbances when worn. No other ACUVUE® Brand of Contact Lenses are included in this voluntary recall.

2. Lazard giving rise to the FSCA

Following our rigorous quality inspections, we identified a high density of microbubbles (microscopic voids within the contact lens) observed in specific lots during our quality control checks. These microbubbles won't cause medical harm to your patients but may be associated with visual disturbances such as glare, halo, starburst, and decreased contrast, particularly in low-light conditions. If patients experience these related visual issues, they will resolve after the removal of the lens.

2. 3. Probability of problem arising

We have not received any complaints, nor have any adverse events been reported at this time due to this issue.

2. 5. Other information relevant to FSCA

You are receiving this notice because our records indicate that you have received contact lenses impacted by this recall. Below we have shared detailed instructions for what to do with the impacted contact lenses you may have on hand. We regret any inconvenience this may cause and are here to support you by replacing the affected lenses – please contact your local Customer Service Team at <Insert Local Info here> for further information, and to address any questions or concerns about ACUVUE® Brand Contact Lenses.



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	3. Type of Action to mitigate the risk						
3.	1. Action To Be Taken by the User						
	☐ Identify Device ☐ Quarantine Device ☐ Return Device ☐ Destroy Device						
	□ On site device modific	estion / increation					
	☐ On-site device modific	cation / inspection					
	☐ Follow patient manag	ement recommendations					
	☐ Take note of amendment / reinforcement of Instructions For Use (IFU)						
	☐ Other ☐ N	ono					
	□ Ottlei □ N	one					
	1. Review if any of yo	our inventory contains contact len	ses impacted f	rom the lot			
	numbers listed bel		•				
	Brand Name	Product Specification	SKU	Lot			
				Number			
	ACUVUE® OASYS MAX 1- Day MULTIFOCAL	1D MAX MULTIFOCAL 8.4 -2.00 LOW 30P RX	888290919116	J003Q72			
		emove from your inventory all affe		Note: You can			
		other lots not affected by this volu otice on to anyone in your organiz	•	do to bo owere			
		isure that they maintain awarenes					
		s will arrange for you the return ar					
	affected product.	vim arrange for you are retain a	.a ropiacomoni	or arry			
	 Complete the enclosed Customer Reply Form and return via email to Customer 						
		cal Info here> for EVEN IF YOU I					
		ted by this recall. JJVC requires					
	reconciliation purposes with regulatory agencies. The completed Customer						
		l be emailed or faxed within 3 bus	siness days of	receipt of			
	this letter.						
3.	2. Is customer Reply	Required?	Yes				
	(If yes, form attached spec						
3.		ken by the Manufacturer	·				
			odification/inspection				
	☐ Software upgrade	☐ IFU or labelling change					
	☐ Other	□ None					
	Swift measures have been n	ut in place to ensure the issue does	not recur so vou	and your			
	Swift measures have been put in place to ensure the issue does not recur so you and your patients can feel confident you'll continue to receive the high-quality contact lenses you know and						
	trust from ACUVUE®.						

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4. General Information					
4.	1. FSN Type	New			
4.	2. Further advice or information already expected in follow-up FSN?	No			
4.	The Competent (Regulatory) Authority of your country has been informed about this communication to customers. Yes				
4.	4. List of attachments/appendices:	ACUVUE® OASYS MAX 1-Day MULTIFOCAL Lens FSCA CUSTOMER REPLY FORM			
4. 5. Name/Signature		Miranda Angelovski EMEA Commercial Quality Manager			

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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ACUVUE® OASYS MAX 1-Day MULTIFOCAL Lens FSCA CUSTOMER REPLY FORM

Please complete and return immediately <u>EVEN IF YOU HAVE NO STOCK</u> via email: >a href="https://example.com/stock-nc/4">>a href="https://example.com/stock-nc/4">>a href="https://example.com/stock-nc/4">>a href="https://example.com/stock-nc/4">>a href="https://example.com/stock-nc/4">>a href="https://example.com/stock-nc/4">a href="https://example.com/stock-nc/4">>a href="https://example.com/stock-nc/4">a href="https://example.

Please p	olace an "X" in one of the b	ooxes below.				
	We have no stock of le		ne recall. ment:			
	Lot Numb	pers	Quantity of lenses to be Returned			
	Company Name:					
	Address:					
	City, State, Zip Code					
	Country					
	Telephone Number:					
			edges the receipt and understanding			
	of the actions, as stated in the Product Recall letter:					
	Name: (print)					
	Title/Position					
	Signature:					
	Date:					