

FSCA Ref: FCA-134

Date: DD:MMM:YYYY.

Urgent Field Safety Notice Update

Edwards Lifesciences IntraClude[™] Intra-aortic Occlusion Device (Model ICF100) This information updates and replaces information provided in Edwards previous Urgent Field Safety Notice ref: FCA 134 of [May 30, 2019]

For Attention of*:To: <<Customer Name>> <<Customer Address>> <<Customer City, State, Postal Code>> <<Customer Country>>

Contact details of local representative (name, e-mail, telephone, address etc.)* This could be a distributor or local branch of the manufacturer. To be added at the appropriate stage in the different local languages



FSCA Ref: FCA-134

<u>Urgent Field Safety Notice (FSN) Update</u> <u>IntraClude™ Intra-aortic Occlusion Device (Model ICF100)</u> <u>Possibility of Balloon Rupture with the IntraClude™ Intra-aortic</u> <u>Occlusion Device (Model ICF100)</u>

	1. Information on Affected Devices*			
1	1. Device Type(s)*			
	The IntraClude Intra-Aortic Occlusion Device (model number ICF100) is a triple-lumen catheter with an elastomeric balloon near its distal tip designed to occlude the ascending aorta in order to partition the aortic root from arterial circulation. The balloon expands to occlude a range of aorta sizes from 20 to 40 mm. This device is designed to be used in the femoral approach with the Edwards EndoReturn (ER21B or ER23B) arterial cannula or the Edwards Introducer Sheath (IS19A). The shaft is provided with an extended strain relief, which tapers from 10.5 Fr to the remaining 9 Fr catheter, and is designed to prevent kinking at the hub and to avoid compressing the shaft when the hemostasis valve of the Edwards EndoReturn (ER21B or ER23B) arterial cannula or Edwards Introducer Sheath (IS19A) is firmly closed. The large central lumen of the IntraClude Device serves three functions: accommodating the guidewire, delivering cardioplegia solution to the aortic root, and venting fluid and air from the aortic root pressure monitoring. The hub has two flexible extension tubes with an integrated luer connection to provide access for the accessories. The shaft is provided with markers to indicate the insertion depth. A blue Clamp-Lock device, provided on the extended strain relief portion, allows the IntraClude Device to be locked in position. The IntraClude Intra-Aortic Occlusion Device is indicated for use in patients undergoing cardiopulmonary bypass. The IntraClude Intra-Aortic Occlusion Device sculues and vents the ascending aorta when the balloon is inflated. The device's central lumen allows delivery of cardioplegia to arrest the heart. The pressure lumen allows monitoring of the aortic root pressure.			
1	2. Commercial name(s)			
	IntraClude Intra-Aortic Occlusion Device			
	3. Unique Device Identifier(s) (UDI-DI) Complete when this becomes available.			
1	4. Primary clinical purpose of device(s)*			
•	The IntraClude intra-aortic occlusion device is indicated for use in patients undergoing cardiopulmonary bypass. The IntraClude intra-aortic occlusion device occludes and vents the ascending aorta when the balloon is inflated. The device's central lumen allows delivery of cardioplegia to arrest the heart. The pressure lumen allows monitoring of the aortic root pressure.			
1	5. Device Model/Catalogue/part number(s)*			
•	ICF100			



FSCA Ref: FCA-134

1	6. Software version
	N/A
1	7. Affected serial or lot number range
-	All lot numbers of Edwards Lifesciences IntraClude™ intra-aortic occlusion device
	(Model ICF100)
1	8. Associated devices
	N/A

	2 Reason for Field Safety Corrective Action (FSCA)*			
2	1. Description of the product problem*			
-	If the IntraClude balloon bursts during cardiopulmonary bypass, the heart can fill and warm, the operative site may be obscured, and the device will need to be exchanged or operative strategy would need to change, including placement of an external cross-clamp, conversion to an open procedure, or performing the procedure under fibrillation. It is possible for an injury to occur as a result of a balloon burst that leads to a change in operative strategy.			
2	2. Hazard giving rise to the FSCA*			
	The issue involves balloon bursts during IntraClude use. The IntraClude device is essential to occlude the aorta and provide the necessary cardiac isolation required to perform minimally invasive cardiac surgery procedures. If the IntraClude balloon bursts during cardiopulmonary bypass, the heart can fill and warm, the operative site may be obscured, and the device will need to be exchanged or operative strategy would need to change, including placement of an external cross-clamp, conversion to an open procedure, or performing the procedure under fibrillation			
2	3. Probability of problem arising			
	Harm may occasionally occur			
2	4. Predicted risk to patient/users			
	The risk is considered as High			
2	5. Further information to help characterise the problem			
	N/A			
2	6. Background on Issue			
	Edwards received some complaints regarding Edwards IntraClude Intra-Aortic Occlusion Device (Model ICF100) related to balloon rupture/puncture, the complaint rate is within the expected occurrence rate, however there were an increase of the number of complaints received between January and April 2019. The issue involves balloon bursts during IntraClude use.			
2	7. Other information relevant to FSCA			
	N/A			



FSCA Ref: FCA-134

	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 modification/inspection					
		Follow patient management recommendations					
		\Box Take note of amendment/reinforcement of Instructions For Use (IFU)					
		□ Other □ None					
		Review this field safety notice to understand the potential safety risk a) Complete and return the Customer Acknowledgment Form. b) Contact Customer Service to arrange return and replacement of affected devices, and					
		c) Return affected devices to Edwards with the Return Goods Authorization (RGA) provided					
		Distribute this notice within your organization or to any organization where the potentially affected devices have been transferred. If you have further distributed this product, notify your customers to the user level. Report any balloon failures to Edwards Lifesciences.					
3.	2	By when should the Within five (5) business days of receiving this					
		action be completed? notice to Customer Service					
3.	3.	Particular considerations for: Choose an item.					
		Is follow-up of patients or review of patients' previous results recommended? Choose an item.					
		Section not applicable					
3.		Is customer Reply Required? * Yes					
0	•	yes, form attached specifying deadline for return)					
3.	ວ.	Action Being Taken by the Manufacturer					
		☑ Product Removal □ On-site device modification/inspection					
		□ Software upgrade □ IFU or labelling change					
		□ Other □ None					
		N/A					
3	6.	By when should the By 27-Sep-2019 action be completed?					



FSCA Ref: FCA-134

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.		



FSCA Ref: FCA-134

	4. General Information*		
4.	1. FSN Type*	Update	
4.	2. For updated FSN, reference number and date of previous FSN	FCA134 – 30 May 2019	
4.	3. For Updated FSN, key new information as follows:		
		nded to all manufacturing lots, so action is required eived and responded to the previous notification.	
4.	 Further advice or information already expected in follow-up FSN? * 		
4	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 representativ		
	a. Company Name b. Address	Edwards Lifesciences LLC	
	c. Website address	One Edwards Way Irvine, CA 92614 www.edwards.com	
4.	8. The Competent (Regulatory) Authority of your country has been informed about th communication to customers. *		
4.	9. List of attachments/appendices:	Customer Reply Form	
4.	10. Name/Signature	Frédérique Pedretti Vice President, Quality Edwards Lifescience	

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.



FSCA Ref: FCA-134

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..*

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



Customer Reply Form

1. Field Safety Notice (FSN) information	
FSN Reference number*	FCA134
FSN Date*	Pre-filled by manufacturer
Product/ Device name*	Edwards Lifesciences IntraClude™ Intra- aortic Occlusion Device (Model ICF100)
Product Code(s)	ICF100
Batch/Serial Number (s)	All lot numbers of Edwards Lifesciences IntraClude™ intra-aortic occlusion device (Model ICF100)

2. Customer Details	
Account Number	
Healthcare Organisation Name*	
Organisation Address*	
Department/Unit	
Shipping address if different to above	
Contact Name*	
Title or Function	
Telephone number*	
Email*	

3. C	3. Customer action undertaken on behalf of Healthcare Organisation				
	I confirm receipt of the Field Safety Notice and that I read and understood its content.	Customer to complete or enter N/A			
	I performed all actions requested by the FSN.	Customer to complete or enter N/A			
	The information and required actions have been brought to the attention of all relevant users and executed.	Customer to complete or enter N/A			
	I have returned affected devices - enter number	Qty: Lot/Serial Number: Date Returned (DD/MM/YY):			
		Qty:	Lot/Serial Number:	Date Returned(DD/MM/YY):	



	of devices returned and date complete.	N/A	Comments:
	I have destroyed affected devices – enter number	Qty:	Lot/Serial Number:
	destroyed and date	Qty	Lot/Serial Number:
	complete.	N/A	Comments:
	No affected devices are available for return/ destruction	Customer to complete or enter N/A	
	Other Action (Define):		
	I do not have any affected devices.	Customer to complete or enter N/A	
	I have a query please contact me (e.g. need for replacement of the product).	Customer to enter contact details if different from above and brief description of query	
Print Name*		Customer print name here	
Signature*		Customer sign here	
Date*			

4. Return acknowledgement to sender	
Email	Pre-filled by manufacturer/sender/requester
Customer Helpline	Pre-filled by manufacturer/sender/requester
Postal Address	Pre-filled by manufacturer/sender/requester
Web Portal	Pre-filled by manufacturer/sender/requester
Fax	Pre-filled by manufacturer/sender/requester
Deadline for returning the customer reply	Pre-filled by manufacturer/sender/requester
form*	

Mandatory fields are marked with *

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.