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FSN Ref: INTCOMP131-FSN

FSCA Ref: INTCOMP131-FSCA

Date: DD MM YYYY

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.

Urgent Field Safety Notice Rocket KCH™ Fetal Bladder Drain R57405 Device Destruction

For Attention of: Persons responsible for medical device vigilance / risk management

Clinicians in the fetal medicine department

Distributors of the device

Contact details of local representative:

For further information, please contact: Regulatoryaffairs@rocketmedical.com

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Urgent Field Safety Notice Rocket KCH™ Fetal Bladder Drain R57405 Device Destruction Material Non-Conformance

I. Information on Affected Devices I. Device Type(s) Rocket KCH™ Fetal Bladder Drain Procedure kit is a sterile, single-use device intended to create a fetal-amniotic shunt to treat fetal lower urinary tract outflow obstruction by allowing the urine to flow from the baby's bladder into the amniotic sac, bypassing the urinary tract. The device contains a double pigtail stent with an outer tube diameter of 2.1mm and inner tube diameter of 1.5mm. Fetal coil Maternal coi 2. Commercial name(s) Rocket KCH™ Fetal Bladder Drain Rocket KCH™ Fetal Bladder Catheter 3. Unique Device Identifier(s) (UDI-DI) ı R57405 4. Primary clinical purpose of device(s) The device is indicated for use in fetal bladder decompression following the diagnosis of fetal post-vesicular obstructive uropathy in fetuses of 18-32 weeks gestation. 5. Device Model/Catalogue/part number(s) R57405 6. Software version N/A – This device is not software and nor does it incorporate software.

	2. Reason for Field Safety Corrective Action (FSCA)		
2	I. Description of the product problem		
	An error has been made in which material of an inferior quality was provided and used in the		
	manufacture of the device. It is understood that the difference in the quality of the materials is		
	limited to Quality Controls around their manufacture, the material used in manufacture having		
	lower controls.		
2	2. Hazard giving rise to the FSCA		
	Sales of the device have been suspended whilst we investigate the impact of the use of this		
	material		

7. Affected serial or lot number range

N/A – There are no other devices associated with this FSN.

00000000466788, 000000000472169. **8. Associated devices**

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2	3. Probability of problem arising			
	Further	urther evaluation is required. To date, no incidents have been reported as a consequence of this		
	issue.			
2	4.	Predicted risk to patient/users		
	It is no	t possible to estimate the risk to patients until further evaluation of this issue has been		
	comple	ted.		
2	5. Further information to help characterise the problem			
	N/A – 1	No further information.		
2	6. Background on Issue No incidents have been reported as a consequence of this issue.			
	An error has been made in which material of an inferior quality was used in the manufacture of			
	the dev	ice. It is understood that the difference in the quality of the materials is limited to Quality		
	Controls around their manufacture, the material used in manufacture having lower controls. We			
	do not know the impact of the use of the incorrect material; a review is underway. In the meantime, we have suspended product sales and we are issuing this FSN to address product in			
	the field.			
2	7.	Other information relevant to FSCA		
	Sales o	f the device continue to be suspended. This field safety corrective action is being		
	implem	ented to destroy any unused product on the market. At this time, no action is considered		
	justified	for patients with an implanted device.		

	3. Type of Action to mitigate the risk				
3	I.	Action To Be T	aken by the User		
		☐ Identify Device	☐ Quarantine Device	☐ Return Device	☑ Destroy Device
		☐ On-site device m	odification/inspection		
		☐ Follow patient ma	anagement recommendations		
		☐ Take note of ame	endment/reinforcement of Ins	structions For Use (IFU)	
		□ Other	☐ None		
			entify any KCH™ Fetal Bla tock. Destroy all devices estroyed devices.		
			at you have received this co		•
		has been transferr	aff members are informed red/supplied to another fac ately by providing a copy o	ility or organisation, pl	-
			ased on the information av The device is critical for t		

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implanted with this device. Any potential remedial action, such as replacing the device with an alternative or expediting delivery, is considered to carry greater risk than leaving implanted devices in situ. All queries regarding this FSN should be directed to Rocket Medical PLC through the email address Regulatoryaffairs@rocketmedical.com. 3 2. By when should the Immediately and without delay. action be completed? 3. Particular considerations for: implantable device 3 Is follow-up of patients or review of patients' previous results recommended? Not at this time. Once further testing has been undertaken to help quantify the risk to patients who have had this device implanted, Rocket Medical will issue further advice regarding appropriate follow-up of those patients. 4. Is Customer Reply Required? Yes (Please complete and return applicable form(s).) 3 5. Action Being Taken by the Manufacturer ☑ Product Removal ☐ On-site device modification/inspection ☐ Software upgrade ☐ IFU or labelling change ☐ Other Further actions, including additional testing, are being undertaken to allow return of the device to the market. 3 6. By when should the As soon as possible. action be completed? 7. Is the FSN required to be communicated to the 3 Nο patient /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?

	4. General Information		
4	I. FSN Type	New	
4	2. For updated FSN, reference number and date of previous FSN	N/A – This is a new FSN.	
4	3. For Updated FSN, key new info	rmation as follows:	
	N/A – This is a new FSN.		

N/A.

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Head of Quality and Regulatory Affairs

Rocket Medical PLC



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4	4. Further advice or information	Yes.			
7	already expected in follow-up	1 63.			
	FSN?				
	5. If follow-up FSN expected, what	5. If follow-up FSN expected, what is the further advice expected to relate to:			
4	The follow-up FSN is expected to provide information regarding appropriate follow-up patients previously implanted with this device.				
	6. Anticipated timescale for				
4	follow-up FSN	1 14/ 2021			
	•				
4	7. Manufacturer information				
	(For contact details of local representative re-	fer to page 1 of this FSN)			
	a. Company Name	Rocket Medical PLC			
	b. Address	Sedling Road, Washington, Tyne & Wear, NE38 9BZ, England			
	c. Website address	www.rocketmedical.com			
4	8. The Competent (Regulatory) A	uthority of your country has been informed			
	about this communication to cus				
4	9. List of attachments/appendices:	- Customer Response Form			
4.	10. Name/Signature				
		Ruth Sharples			

Transmission of this Field Safety Notice		
This notice needs to be passed on to 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 of 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 Response Form

I. Field Safety Notice (FSN) information	
FSN Reference number	INTCOMP131-FSN
FSN Date	DD MM YYYY
Product/ Device name	Rocket KCH™ Fetal Bladder Drain
Product Code(s)	R57405
Batch/Serial Number (s)	00000000466788, 00000000472169.

2. Customer Details	
Healthcare Organisation Name	
Organisation Address	
Department/Unit	
Contact Name	
Title or Function	
Telephone number	
Email	

3. C	ustomer action undertak	en on behalf of Healthcare Organisation
	I confirm receipt of the Field Safety Notice and that I read and understood its content.	Comment
	I have/will perform all actions requested by the FSN.	Comment
	The information and required actions have been brought to the attention of all relevant users.	Comment
	I have destroyed the following number of devices:	Number of devices:
	The Batch/Serial Number (SN or LOT) for devices destroyed are:	Serial / LOT number (required for replacement / reimbursement):
	I do not have any affected devices.	Comment
Print I		
Signat	ure	
Date		

4. Return acknowledgement to:	
Email	regulatoryaffairs@rocketmedical.com
Subject of e-mail	"INTCOMP131-FSN Response"

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Deadline for returning the Customer Response	Immediately / As soon as possible.
form	