

FSCA Ref: PR6



Date: 26/05/203

## Urgent Field Safety Notice Device Commercial Name

For Attention of: all affected distributors and users

Contact details of the manufacturer. Altomed Ltd, 2 Witney Way, Boldon Business Park, Boldon, Tyne and Wear, NE35 9PE, United Kingdom. Quality & Regulatory Manager: Bethany Garside E-Mail: Bethany.garside@altomed.com Tel: +44 (0) 191 519 0111

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

	1. Information on Affected Devices*
1.	1. Device Type(s)*
	Damato Ruthenium Plaque Template - A sterile device in the form of a dome, designed to be
	placed over a tumour that is inside the eye to help determine optimal positioning of an eye
	brachytherapy plaque.
1.	2. Commercial name(s)
	Damato Ruthenium Plaque Template
1.	3. Unique Device Identifier(s) (UDI-DI)
	05055505156900,
	05055505156894,
	05055505156887
1.	<ol> <li>Primary clinical purpose of device(s)*</li> </ol>
	A sterile device in the form of a dome, designed to be placed over a tumour that is inside the
	eye to help determine optimal positioning of an eye brachytherapy plaque.
1.	<ol><li>Device Model/Catalogue/part number(s)*</li></ol>
	A7075CIB, A7075CIA, A7075COC
1.	6. Software version
	N/A
1.	7. Affected serial or lot number range
	A7075CIB = 01108, 01301, 01508, 01300. A7075CIA = 01508, 01108, 01300, 01301. A7075COC =
	01108, 01107.
1.	8. Associated devices
	N/A

	2 Reason for Field Safety Corrective Action (FSCA)*			
2.	<ol> <li>Description of the product problem*</li> </ol>			
	Our international distributor informed us of a complaint they received from one of their			
	customers. The CIB template (REF A7075CIB, LOT 01108) used in surgery did not precisely match			
	up with the suture holes of the related CIB ruthenium plaque supplied. On further investigation it			
	was found that the suture holes on the related ruthenium plaques also do not precisely align with			
	template variants A7075CIA and A7075COC.			
2.	2. Hazard giving rise to the FSCA*			
	No direct safety issue. Potential for extended surgery time if the related plaque suture holes do			
	not precisely align with the sutures placed using the template.			
2.	3. Probability of problem arising			
	Assessed as low given that multiple surgeries (estimated less than 200) may have been performed			
	without any reported incident. However, given the potential for extension of surgery time all lot			
	numbers of all three products are being withdrawn as a precaution in order that the basis for the			
	mismatch described can be further investigated and addressed.			
2.	<ol> <li>Predicted risk to patient/users</li> </ol>			
	Negligible – extended surgical intervention.			
2.	5. Further information to help characterise the problem			

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	n/a		
2.	6. Background on Issue		
	Altomed were made aware of this issue when our international distributor highlighted a customer		
	complaint that the holes of the template did not fit the holes of the CIB-plaque. This resulted in a		
	two-hour prolonged surgery for one patient, with revised suture holes and extra exposure for		
	patient and personnel. The root cause of the error is not fully known yet, but likely relates to a		
	design specification mismatch between the template dimensions and the dimensions of the		
	related ruthenium plaques with which they are used. Therefore, we are presuming at this stage		
	that all lot numbers are affected.		
2.	7. Other information relevant to FSCA		
	n/a		

	3. Type of Action to mitigate the risk*				
3.	1.	1. Action To Be Taken by the User*			
		⊠ Identify Device ⊠ Quar	antine Device	🛛 Return Device	Destroy Device
		□ On-site device modification	n/inspection		
			in inspection		
		□ Follow patient manageme	ent recommendations	S	
		□ Take note of amendment,	reinforcement of Ins	structions For Use (	IFU)
		□ Other □ None			
	Ret	curn devices to Altomed. Repl	acements or credit w	vill be issued.	
		•			
3.	2.	By when should the action be completed?	As soo	n as possible	
		be completed!			
3.	3.	Particular considerations for	: Choose ai	n item.	
		la fallano un af nationte an va	ious of notionts' and		
		Is follow-up of patients or review of patients' previous results recommended? Yes			imended?
		If any of the affected devices have been used, the attending surgeon should be consulted			
2	4	for an assessment of whether		ent may have been	
3.		4. Is customer Reply Required? * Yes (If yes, form attached specifying deadline for return)			
3.		5. Action Being Taken by the Manufacturer			
		0 /			
		⊠ Product Removal	] On-site device mod	ification/inspection	ı
		□ Software upgrade □	] IFU or labelling cha	nge	
		□ Other □	None		
		Provide further details of the action(s) identified.			



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3	6.	By when should the action	As soon as possible	
		be completed?		
3.	7.	Is the FSN required to be communicated to the patient /lay No		No
		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 a	in item.	

	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 informatio	n 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	e further advice expected to relate to:
4		n/a	
	6.	Anticipated timescale for follow-up	n/a
4		FSN	
4.	7.	Manufacturer information	
	(Fo	or contact details of local representative	
		a. Company Name	Altomed Limited
		b. Address	2 Witney Way, Boldon Business Park, Tyne and Wear. NE35 9PE
		c. Website address	www.altomed.com
4.	8.	The Competent (Regulatory) Author	ity of your country has been informed about this
		communication to customers. *	
4.	9.	List of attachments/appendices:	PR6 FSN Customer Reply Form/ Distributor Reply
4	10		Form
4.	10	Name/Signature	Bethany Garside
			QA/RA 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..\*

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