## **Urgent Field Safety Notice: RA2023-3471895**

### December XX, 2023

#### **Affected product**

<b>Product Family Names:</b>	Scorpio, Duracon, PCA, Trident
Identification of the	See Part/Lot Number Attachment: PFA RA2023-3471895 starting on page 3
Affected Products:	

Dear Customer,

Stryker has initiated a voluntary, lot specific recall for the devices listed in the Part/Lot Number Attachment: PFA RA2023-3471895. The intent of this letter is to list all known hazards and harms potentially associated with the below noted issue and list the risk mitigation factors associated with the use of the product.

#### Issue

The industry guidance for storage duration of Ultra High Molecular Weight Polyethylene (UHMWPE) raw material used in the manufacture of soft implant bearing/articulating material of prosthetic joints is less than 5 years. Product manufactured using UHMWPE raw material over 5 years of age has the potential for elevated levels of oxidation. Oxidation within UHMWPE can have an impact on its material properties.

Stryker has become aware that the devices listed in the Part/Lot Number Attachment: PFA RA2023-3471895 have been manufactured with UHMWPE raw material aged over 5 years.

#### **Potential Hazards**

- Excessive Wear Debris
- Material Fragments
- Fractured Device

#### **Potential Harms**

- Revision Surgery
- Pain
- Inflammation

#### **Risk Mitigations**

Product with a high oxidation index can become discolored. However, not all product with a high oxidation index becomes discolored. In instances where the product is discolored, the issue may be recognized by the user.

#### **Recommendations for patients already implanted with an impacted device**

Post-market and National Registry Joint data were evaluated for devices in scope. Stryker identified no trends for the potential hazards. Patients treated with an impacted product identified should continue to be followed per the normal protocol established by his or her surgeon(s). There are no recommended changes to the frequency of the standard follow-up care protocol. Additional or more frequent patient monitoring or follow-up may be required in accordance with clinical judgment.



### **Actions needed**

Our records indicate that you may have received the affected product(s). It is Stryker's responsibility as the manufacturer to ensure that customers who may have received these affected products also receive this important communication. We therefore request that you read this notice carefully and complete the following actions.

- 1. Immediately check your internal inventory and quarantine all subject devices pending return to Stryker.
- 2. Circulate this Field Safety Notice internally to all interested/affected parties.
- 3. Maintain awareness of this notice internally until all required actions have been completed within your facility.
- 4. Discontinue use of the recalled devices identified in the affected product list (see Part/Lot Number Attachment: PFA RA2023-3471895 starting on page 3).
- 5. Inform Stryker if any of the subject devices have been distributed to other organizations.
  - a. Please provide contact details so that Stryker can inform the recipients appropriately.
  - b. If you are a Distributor, note that you are responsible for notifying your affected customers.
- 6. Please inform Stryker of any adverse events concerning the use of the subject devices.
  - Please comply with any local laws or regulations concerning the notification of adverse a. events to your National Competent Authority.
- 7. Complete the attached customer response form. It may be that you no longer have any physical inventory on site. Completing this form will allow us to update our records and will also negate the need for us to send any further unnecessary communications on this matter. Therefore, please complete even if you no longer have any of the subject devices in your physical inventory.
- 8. Return the completed form to your nominated Stryker Representative (indicated below) for this PFA.
  - a. On receipt of the form, a Stryker Representative will contact you to organize any applicable ongoing actions.

We request that you respond to this notice within 7 calendar days from the date of receipt.

Please respond event if you do not have a record of receiving affected inventory. This will enable us to update our records and negate the need to send unnecessary reminder letters. Your timely response will enable us to update our records and negate the need to send reminder notices.

Your designated contact person for this action is given below. Should you have any queries concerning

this matter please do not hesitate to contact them directly.

Name:

*Position: email:* 

On behalf of Stryker, we thank you sincerely for your help and support in completing this action within the target date and regret any inconvenience that may be caused. We would like to reassure you that Stryker is committed to ensuring that only conforming devices, meeting our high internal quality standards, remain on the market.

Sincerely,

Part/Lot Number Attachment: PFA RA2023-3471895						
Part Number	Product Description	Lot Numbers			GTIN	
3044-0026	SCORPIO RECESSED PATELLA	PVDY	V644	V355	07613327033779	
		TEN1	MHHE	55LK		
		AMN1	4T1K	JR6R		
3044-0028	SCORPIO RECESSED PATELLA	A59N	EP7J	X79L	07613327033793	
3044-0030	SCORPIO RECESSED PATELLA	PD99	6AHH	RY3J	07613327033809	
3044-0032	SCORPIO RECESSED PATELLA	6LR4	R14V	N882	07613327033823	
3052-0515	SERIES II TIBIAL BEAR INSERT	WK4PX0			07613327034042	
3052-0524	SERIES II TIBIAL BEAR INSERT	К349ТР			07613327034073	
72-4-0321	SCORPIO TS TIB INSERT	TA21X9			07613327034349	
72-4-0510	SCORPIO TS TIB INSERT	H43RDK	768NEY		07613327034363	
		WR1RE3	4J13P1			
72-4-7512	SCORPIO TS TIB INSERT	5M6LM1			07613327034790	
72-4-7514	SCORPIO TS TIB INSERT	482LM8	HH43K7		07613327034820	
72-4-7516	SCORPIO TS TIB INSERT	TN1746	KK82WY		07613327034806	
72-4-7518	SCORPIO TS TIB INSERT	9P8360	AK8393		07613327034813	
72-4-7521	SCORPIO TS TIB INSERT	VA0X4T			07613327034837	
72-4-7524	SCORPIO TS TIB INSERT	V68WW1	93400601		07613327034844	
72-15-0324	Scorpio-Flex Ttl Kn P-S Tib Brg Insrt Asy	53867101			07613154020133	
72-16-0908	Scorpio-Flex Ttl Kn CR Tib Brg Insrt Asy	42968601			07613154020638	
72-16-0910	Scorpio-Flex Ttl Kn CR Tib Brg Insrt Asy	43103101			07613154020645	
73-0110	SCORPIO TOTAL KNEE MEDIALIZED	79E0	WEKJ	33ND	07613327050288	
	DOME PATELLA-SZ #11	274P	LH82			
73-0510	SCORPIO TOTAL KNEE MEDIALIZED	57A2	E10K	WJ3L	07613327050301	
	DOME PATELLA-SZ #5	3WP6	X49N	38TY		
73-0710	SCORPIO TOTAL KNEE MEDIALIZED	RLAR	Y373	6LN7	07613327050295	
	DOME PATELLA-SZ #7	2P84	58LA	WEL3		
		EW6V	VXAL	NDK3		
		90AV	4201	25WJ		
		4JHN	A2AN	T2JW		
		65A3	5YP8	YVWT		
		22MP	X0P9	7DR0		
		NNE7	9A8T	DMVL		
73-0910	SCORPIO TOTAL KNEE MEDIALIZED	X1WP	0TNX	J72T	07613327050332	
	DOME PATELLA-SZ #9	6KE9	WN5E	M73H	0,01002,000002	
		DL4V	2EM2	HM8H		
		HHPW	96N0	MEM7		
		1NL2	HW7E	M8HN		
		4JHW	T9NM	WL3Y		
73-2110	SCORPIO TOTAL KNEE CONCENTRIC DOME PATELLA-SZ #11	LT28	4TXX		07613327033816	
73-2710	SCORPIO TOTAL KNEE CONCENTRIC	V25H	5MYL	845P	07613327033847	
	DOME PATELLA-SZ #7	MD9E	Н9Т8			
73-2910	SCORPIO TOTAL KNEE CONCENTRIC DOME PATELLA-SZ #9	A02W	75TX		07613327033854	

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Part Number	Product Description		Lot Numb	oers	GTIN
		NPPV	6RLN		
73-3110	SCORPIO U-DOME PATELLA	XEA7	4902	76KN	07613327033861
		8P83	YNP7	K4P9	
		690Y	3AYN	792H	
		MRM3	AYHM	43T8	
		67D4	W3MD	MW40	
73-3308	SCORPIO U-DOME PATELLA	2H58	8H52	X0H9	07613327033878
		2J0W	64MJ	70E9	
73-3508	SCORPIO U-DOME PATELLA	ЕРХН	N8P6	D06K	0761332703388
		DYAE	T355	01JR	
		XVK2	8HDN	JPA4	
		PVP9	A3EV	0AYL	
		LJOK	5LKW	W424	
		12R3	R04Y	53T1	
			RKT1	Y7X2	
73-3708	SCORPIO U-DOME PATELLA	RDWN	TXL1	8PWK	07613327033892
10 0100		4W3T	53RN	R9A4	0,01002,0000,
		V768	KR24	NWXA	
		VJ05	M3T4	D47L	
		TWT6	VE90	5JVL	
		ADK0	EKK2	D16P	
		855	36L1	E2LP	
		D6D9	NH6E	927H	
		0007	NIIOL	JR5N	
73-3710	SCORPIO U-DOME PATELLA	RNT5	5A2R	EOVH	07613327033908
/3-3/10	SCORFIG G-DOME FATELLA	3YNP	98AH	3PA0	0/01332/033900
		PEA1	T1JH	05L4	
73-3910		NNA5	8D72 LEAY	R608 84A4	07(100000000000000000000000000000000000
/3-3910	SCORPIO U-DOME PATELLA	73JP			0761332703391
		V17W	4NTR	XREK	
		N2MY	R5WA	YNM1	
		PK0D	JN8M	LY5M	
		D3M3	084W	M1AT	
00.0.000		40054504	K6YE	4873	0454654040050
82-2-0908	Scorpio NRG Tibial Brg Insert Assy	42874701	42995601		04546540400734
		42912101			
82-2-0910	Scorpio NRG Tibial Brg Insert Assy	42912201	42961901		04546540400742
620-00-28J	HOWMEDICA OSTEONICS TRIDENT 0 deg POLY INSERT	38059601			Inactive
620-00-32J	HOWMEDICA OSTEONICS TRIDENT 0 deg POLY INSERT	29518001			Inactive
6302-6-107	P7 28MM 10 DEGREE +4MM INSERT	61311701	62122001		0761315307615

Part/Lot Number Attachment: PFA RA2023-3471895				
Part Number	Product Description	Lot Numbers	GTIN	
6302-6-307	P7 32MM 10 DEGREE +4MM INSERT	62112001 62120801	07613153076407	
6637-0-028	LEFT SMALL-PCA MOD.REV.DUR.REV.INSERT	42771401	04546540305831	
6637-0-228	LEFT LARGE-PCA MOD.REV.DUR.REV.INSERT	54537601	04546540306012	
6637-0-231	LEFT LARGE-PCA MOD.REV.DUR.REV.INSERT	42787901	04546540306029	
6637-0-328	LEFT X-LARGE-PCA MOD.REV.DUR.REV.INSERT	42579701	04546540306104	
6637-0-631	LEFT SMALL-PCA MOD.REV.DUR.REV.INSERT	36272501 42380501 42771601	04546540306203	
6637-0-831	RGT.LARGE-PCA MOD.REV DUR.REV.INSERT	47475901	04546540306388	
6637-0-928	RGT.LARGE-PCA MOD.REV DUR.REV.INSERT	41875401	04546540306463	
6637-0-931	RGT.LARGE-PCA MOD.REV DUR.REV.INSERT	57403601	04546540306470	
6637-4-031	PRIMARY REV.TIB.INSERT-DURACON	36273601 36609701	04546540306920	
6637-4-231	"LARGE PRIMARY REV.TIB.INSERT- DURACON	33164901	04546540307101	
6642-1-709	DURATION A-P LIPPED TIB.INSERT- DURAC	52098201 56002801 50677001 54184201 56662001	04546540318145	
6642-1-911	DURATION A-P LIPPED TIB.INSERT- DURAC	56000801 57491601	04546540318299	
6642-2-200	DURATION PLASTIC PATELLA-DURACON	383541	04546540318398	
6728-2-609	DUR PCA MTK REV INS LFT	50884601	04546540322876	
6728-2-611	DUR PCA MTK REV INS LFT	52814401	04546540322883	
6728-2-709	DUR PCA MTK REV INS RT	51828501	04546540322944	
6728-2-711	DUR PCA MTK REV INS RT	52177701	04546540322951	
6742-1-411	PS LIPPED TIBIAL INSERT ASSY DURACON	584223	04546540324108	
6742-1-413	PS LIPPED TIBIAL INSERT ASSY DURACON	559469 584958 561926	04546540324115	
6742-1-416	PS LIPPED TIBIAL INSERT ASSY DURACON	571630 584840 582549	04546540324122	
7291-0324	TIBIAL BEARING INSERT SERIES P-S I ASSY	59065701	04546540117144	

### **Business Reply Form - response required**

### **Urgent Field Safety Notice: RA2023-3471895**

### December XX, 2023

<b>Product Family Names:</b>	Scorpio, Duracon, PCA, Trident
Identification of the Affected Products:	See Part/Lot Number Attachment: PFA RA2023-3471895 starting on page 3

I have received the **Urgent Field Safety Notice** letter from Stryker dated December XX, 2023, stating that the company has initiated a voluntary recall on the above referenced affected products.

Please complete the form even if you do not have inventory. This will preclude us from following up.

Customer information			
Customer name:			
Name of person completing this form:			Title:
Direct phone number:		_Email	
Address:			City:
Postal code:	_Country:		

If affected inventory, please provide the information below. Attach additional sheet if needed.

Product code	Lot number	Qty quarantined	Qty destroyed	Qty returned

### We have not located any of these devices in our inventory (please add check mark to box):

If you have further distributed subject devices, please provide information below:

Facility Name	Facility Address	Contact person	Product code	Lot number	Qty

I have read and understand the instructions provided and acknowledge receipt of the subjected Field Safety Notice. I also agree to further distribute and communicate this important information from this letter to those whom I have distributed any of subjected devices noted in this letter.

Name (print): \_\_\_\_\_\_ Signature: \_\_\_\_\_ Date: \_\_\_\_\_

PLEASE COMPLETE THIS FORM WITHIN 7 CALENDAR DAYS AND RETURN IT BY USING THE EMAIL OR FAX\_\_\_\_\_