

24 January, 2018

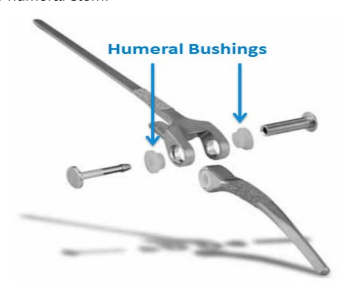
To: Risk Managers

Subject: URGENT MEDICAL DEVICE RECALL

Affected Product: Coonrad/Morrey Elbow Humeral Stem

Item Number	Lot Number	UDI Number
32-8105-027-04	63694912	(01)0088902427 4006(17)220731(10)63694912

Zimmer Biomet is conducting a lot specific medical device recall for the Coonrad/Morrey Humeral Stem. A field complaint investigation determined that the humeral bushing components may not have been packaged with the humeral stem.



	Risks	
Describe immediate health	Most Probable	Highest Severity
consequences (injuries or illness) that may result from use of or exposure to the product issue.	Delay of Surgery (<30 Minutes)	Delay of Surgery (<30 Minutes)
Describe long range health	Most Probable	Highest Severity
consequences (injuries or illness) that may result from use of or exposure to the product issue.	None	None

Our records indicate that you may have received one or more of the affected products. The affected units were distributed in July 2017 and August 2017.



Risk Manager Responsibilities:

- 1. Review this notification and ensure that affected personnel are aware of the contents.
- 2. If you have affected product at your facility, assist your Zimmer Biomet sales representative and quarantine all affected product. Your Zimmer Biomet sales representative will remove the affected product from your facility.
- Complete Attachment 1 Certificate of Acknowledgement and send to <u>CorporateQuality.PostMarket@zimmerbiomet.com</u>. This form must be returned even if you do not have affected products at your facility.
- 4. Retain a copy of the acknowledgement form with your recall records in the event of a compliance audit of your facility's documentation.
- 5. If you have further questions or concerns after reviewing this notice, please call customer service at 574-371-3071 between 8:00 am and 5:00pm EST, Monday through Friday. Calls received outside of call center operating hours will receive a voicemail prompt or be transferred to an on-call representative in the event of an emergency. Alternatively, your questions may be emailed to CorporateQuality.PostMarket@zimmerbiomet.com.

Other Information

This medical device recall was reported to the U.S. Food and Drug Administration and will be reported to other Competent Authorities, Notified Bodies, and Regulatory Authorities as required.

- Med Watch Reporting: Adverse reactions or quality problems experienced with the use of this product may be reported to the FDA's Med Watch Adverse Event Reporting program either online, by mail, or by fax.
- Online: www.fda.gov/medwatch/report.htm
- Mail: Use postage paid, pre-addressed form FDA 3500, available at: www.fda.gov/MedWatch/getforms.htm
- Fax: 1-800-FDA-0178

Under 21 CFR 803, manufacturers are also required to report any serious injuries where a product has contributed or may have contributed to the event. Please keep Zimmer Biomet informed of any adverse events associated with this product or any other Zimmer Biomet product by emailing Zimmer.PER@zimmerbiomet.com.

Thank you for your assistance. We regret any inconvenience caused by this recall.

Sincerely,



Post Market Surveillance & Regulatory Compliance Director



ATTACHMENT 1Certificate of Acknowledgement

<u>IMMEDIATE RESPONSE REQUIRED – TIME SENSITIVE ACTION NEEDED</u>

	uct: CM Humeral Stem	rieiu	Action Reference: ZFA2017-0008
	Please check or	ne as a _l	pplicable:
	☐ Hospital Facility		☐ Surgeon
	Do you have affected p (Hospital Facility Only: Please		
	Yes, we currently have one	or more	affected items in our facility.
	☐ No, we currently have no af	fected ite	ems in our facility.
I notice.			
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