

# To the attention of Medical Device Vigilance responsible / Central Pharmacy

Saint Priest, 23 September 2024

# URGENT - FIELD SAFETY NOTICE - Codman® Surgical Patties & Surgical Strips - RECALL

## Legal manufacturer:

INTEGRA LIFESCIENCES PRODUCTION CORPORATION - 11 Cabot Boulevard - Mansfield, MA 02048 USA - SRN: US-MF-000009189

Swiss Representative: INTEGRA LIFESCIENCES SERVICES (Switzerland) LTD – Fidulem SA, Avenue Mon-Repos 24 – 1005 – Lausanne – Suisse – CHRN-AR: 20001538

#### Medical device:

Codman® Surgical Patties & Strips are manufactured of COTTONOID® Material with x-ray detectable markers. All patties have a suture string attached for ease in performing postsurgical count verification.

## Primary clinical purpose of device:

The surgical patties and surgical strips are indicated for the use in protection of tissue, including brain and other tissues of the central nervous system, during surgery.

#### Concerned references:

PATTIES			S	STRIPS		
801396	801401	801407	801449	801454		
801397	801402	801408	801450	901455		
801398	801403	801409	801451	801456		
801399	801404		801452	801457		
801400	801406		801453			



Dear Valued Integra Customer,

Integra LifeSciences is voluntarily issuing this Field Safety Notice for the recall of Codman® Surgical Patties & Surgical Strips products listed in Table 1 below.

During an internal investigation, Integra LifeSciences identified higher-than-expected levels of endotoxin within the raw material used to produce Codman Surgical Patties and Strips that may have resulted in out-of-specification levels of endotoxin in those finished goods. Consequently, while the endotoxin levels identified were higher than expected, the possibility of adverse health consequences actually occurring remains remote (see Risk to Health below).

#### Risk to health

Per the Health Hazard Evaluation conducted for this issue, adverse health consequences resulting from higher-than-expected levels of endotoxins may include mild febrile response, and/or mild local transitory inflammation, hypotension, or nausea.

If you have already used the products affected by this recall and standard operative care was followed, there is no additional patient follow-up required.

<u>Please note that there have been zero (0) complaints received relating to the potential harms identified</u> in the "Risk to Health" section.

**Table 1: Product Information** 

Manufacturer' s Product Number (Catalog #)	Product Name (Description)	UDI Number	Lot Number	Expiration Date	Distribution Dates (DD/MM/YYYY)
801396	CODMAN MICR PATIE RND/200	20886704036446	All lot numbers distributed	unexpired	All lots distributed
801397	SURGPAT X-RAY1/4X11/2-200		between 01- Aug-2019 to 31-	lots	between 01/08/2019 to
801398	SURG PAT XRAY 1/4X3 -200	10381780514947, 20886704036460	July-2024		31/07/2024
801399	SURG PATXRAY 1/4X1/4-200	10381780514954, 20886704036477			
801400	SURG PATXRAY 1/2X1/2-200	10381780514961, 20886704036484			
801401	SURG PATXRAY 3/4X3/4-200	10381780514978, 20886704036491			
801402	SURG PAT XRAY 1/2X1 -200	10381780514985, 20886704036507			
801403	SURG PAT XRAY 1X1 -200	10381780514992, 20886704036514			
801404	SURG PAT XRAY 1/2X1 1/2	10381780515005, 20886704036521			
801406	SURG PAT XRAY 1/2X2 -200	10381780515012, 20886704036538			
801407	SURG PAT XRAY 1/2X3 -200	10381780515029, 20886704036545			
801408	SURG PAT XRAY 1X3 -200	10381780515036, 20886704036552			
801409	SURG PAT XRAY 3X3 -200	10381780515043, 20886704036569			
801449	CODMAN SRG STRP1/8X6-200	10381780515050, 20886704036576			



Manufacturer' s Product Number (Catalog #)	Product Name (Description)	UDI Number	Expiration Date	Distribution Dates (DD/MM/YYYY)
801450	CODMAN SURGSTRIP1/4X6-200	10381780515067, 20886704036583		
801451	CODMAN SURG STRP1/2X6-200	10381780515074, 20886704036590		
801452	CODMAN SURG STRP3/4X6-200	10381780515081, 20886704036606		
801453	CODMAN SURG STRIP1X6-200	10381780515098, 20886704036613		
801454	CODMAN SURGSTRP11/2X6-200	10381780515104, 20886704036620		
801455	CODMAN SURG STRIP2X6-200	10381780515111, 20886704036637		
801456	CODMAN SURG STRIP3X6-200	10381780515128, 20886704036644		
801457	CODMAN SRG STRP31/2X6-200	10381780515135, 20886704036651		

Our records indicate that you may have received products from these lots.

## Actions to be taken by Customers:

- 1. Please review and understand the information provided in this letter.
- 2. If you do have affected units:
  - a. Quarantine the units immediately.
  - b. Check the box on the enclosed form "I do have affected units."
  - c. Record on the table 2. at the bottom of the reply form the total quantity of affected units and lot number(s) that you have.
- 3. If you do not have affected units, check the box, "I do not have affected units."
- 4. Please **return the completed reply form by email to** <u>emea-fsca@integralife.com</u>, or Fax to +33 (0)4.37.47. 59.30. By filling in this form, you confirm that you have received this Safety Notice, and you intend to fully comply with this notification. **We expect a response within 3 weeks.** You also confirm that this notification has been forwarded to every concerned person in your organization.
- 5. At receipt of your form, and if it is noted that you have affected units, Integra Customer Service will contact you and provide a Return Material Authorization (RMA) number and directions to return the affected product(s). A credit note will be processed upon receipt of returned goods (except for consignments).
- 6. We recommend that you retain a copy of the form for your records.

# PLEASE NOTE THAT REGARDLESS OF WHETHER YOU HAVE THE AFFECTED PRODUCT TO RETURN OR NOT – A COMPLETED ACKNOWLEDGEMENT IS REQUIRED

The receipt of this form ensures that Integra has achieved a level of effectiveness in communicating this information.

National Competent Authorities may perform audits of field actions of this nature to verify that our customers have been notified and understand the nature of the field action being taken.

The National Competent Authority of your country has been alerted of this Field Safety Corrective Action.



Thank you for your cooperation with this Field Safety Corrective Action and for returning the attached Reply Form.

Please feel free to contact our Post Market Surveillance Department at <a href="mailto:emea-fsca@integralife.com">emea-fsca@integralife.com</a> for any additional questions. Your cooperation is appreciated, and we thank you for your continued collaboration.

Yours Sincerely,

Post Marketing Surveillance Department

Appendix: Field Safety Notice Reply Form (2 pages)



**Customer Reply Form** 

1. Field Safety Notice (FSN) information	
FSN Reference number	2024-HHE-013
FSN Date	23 September 2024
Product/ Device name	Codman® Surgical Patties & Strips
Product Code(s)	See list in table 2 below
Lots	All lot numbers distributed between
	01/08/2019 to 31/07/2024

3. C	3. Customer action undertaken on behalf of Healthcare Organisation				
S.   C	I confirm receipt of the Field Safety Notice and that I read and understood its content.  I performed all actions requested by the FSN.	or nearmeare Organisation			
	The information and required actions have been brought to the attention of all relevant users and executed.				
	I <u>have</u> affected units - enter number of products and lot number	Fill in the table 2 below			
	I do not have any affected units.				
	I have a query please contact me	Customer to enter contact details if different from above and brief description of query			
Print	Name*	Customer print name here			
Signa	ature*	Customer sign here			
Date*	•				



4. Return acknowledgement to Sender		
Email	emea-fsca@integralife.com	
Customer Helpline	+33 (0) 6 30 20 69 66	
Postal Address	Post Market Surveillance Department Integra Immeuble Séquoia 2, 97 allée Alexandre Borodine Parc technologique de la Porte des Alpes 69800 Saint Priest, France	
Web Portal	https://integralife.eu/	
Deadline for returning the customer reply form*	20/10/2024	

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 action.

**Table 2. List of products** 

Manufacturer's Product Number (Catalog Number)	Product Name (Description)	Lot Number(s) as identified on the box or on the pouch	Quantity (box) Note: partial box counts as a full box
801396	CODMAN MICR PATIE RND/200		
801397	SURGPAT X-RAY1/4X11/2-200		
801398	SURG PAT XRAY 1/4X3 -200		
801399	SURG PATXRAY 1/4X1/4-200		
801400	SURG PATXRAY 1/2X1/2-200		
801401	SURG PATXRAY 3/4X3/4-200		
801402	SURG PAT XRAY 1/2X1 -200		
801403	SURG PAT XRAY 1X1 -200		
801404	SURG PAT XRAY 1/2X1 1/2		
801406	SURG PAT XRAY 1/2X2 -200		
801407	SURG PAT XRAY 1/2X3 -200		
801408	SURG PAT XRAY 1X3 -200		
801409	SURG PAT XRAY 3X3 -200		
801449	CODMAN SRG STRP1/8X6-200		
801450	CODMAN SURGSTRIP1/4X6-200		
801451	CODMAN SURG STRP1/2X6-200		
801452	CODMAN SURG STRP3/4X6-200		
801453	CODMAN SURG STRIP1X6-200		
801454	CODMAN SURGSTRP11/2X6- 200		
801455	CODMAN SURG STRIP2X6-200		
801456	CODMAN SURG STRIP3X6-200		
801457	CODMAN SRG STRP31/2X6-200		