

Smith & Nephew, Inc.
Global Field Actions
1450 Brooks Road
Memphis, TN 38116
Tennessee, USA

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T: 1 800 821 5700 (USA toll free)
www.smith-nephew.com



<Recipients Address>

URGENT FIELD SAFETY NOTICE: Product Recall

Date Issued: 24-November-2025

Reference: R-2025-07

Legal Manufacturer: Smith & Nephew, Inc.

Concerned Devices: BIOSURE HA 6×20 mm / BIOSURE REGENESORB (RG) 5×20 mm
Interference Screws

Product No.	Description	Batch No.	Unique Device Identifier(s)
72201768	BIOSURE HA 6×20 mm interference screw	51315219	03596010611642
72204389	BIOSURE REGENESORB (RG) 5×20 mm interference screw	51306803	00885554036572

Dear Customer:

This letter is to inform you that Smith & Nephew, Inc., has initiated a field action to voluntarily remove one lot of BIOSURE HA 6×20 mm interference screws and one lot of BIOSURE REGENESORB (RG) 5×20 mm interference screws due to a labeling error. Complaints were received indicating a partial mix of packages containing BIOSURE REGENESORB (RG) 5×20 mm interference screws instead of BIOSURE HA 6×20 mm interference screws as described on the product label.

This field action has been reported to the relevant competent authorities.

Patient Impact

Smith+ Nephew recommends that physicians maintain their routine patient follow-up protocol.

Risks to Health	<p>In the most likely scenario, the labeling mismatch is identified during surgery set up, or the screw size difference or presence/absence of fenestrations is identified by the surgeon prior to screw insertion. A backup device is used, and the surgery is completed without a significant delay. There is no hazardous situation and no harm.</p> <p>In the worst case scenario, the discrepancy is not identified, and the incorrect screw is used and inserted. If the screw breaks, there is a remote possibility of a surgical delay greater than 30 minutes to remove the broken screw and use a backup device to complete the surgery.</p>
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<p>Actions to be taken by the user</p>	<ol style="list-style-type: none">1. Ensure that the contents of this Field Safety Notice are read and understood by those within your organisation who may use BIOSURE HA 6×20 mm or BIOSURE REGENESORB (RG) 5×20 mm Interference Screws2. Locate and quarantine affected devices immediately. If you have further distributed the product to other organisations, please inform them at once of this Field Action and provide to them a copy of this letter.3. Please complete the Customer Response form and email or fax it to your national Smith+Nephew agency/distributor.4. Return quarantined product to your national Smith+Nephew agency/distributor.5. Please maintain awareness of this notice and resulting action for an appropriate period to ensure the effectiveness of the corrective action.
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If you or any of the healthcare providers you serve have any questions regarding this information, please contact your national Smith+Nephew agency/distributor.

Smith+Nephew is committed to distribute only products of the highest quality standards and to provide any required support. We regret that this has occurred and any inconvenience it may cause or has caused you, your patients, or your staff.

Thank you for your attention and cooperation.

Customer Response Form

Please read in conjunction with the Field Safety Notice and return the completed and signed Customer Response Form by <date>.

Reference: R-2025-07

Concerned Devices: BIOSURE HA 6×20 mm / BIOSURE REGENESORB (RG) 5×20 mm
 Interference Screws

1. Return Acknowledgement details	
Email	<Local market to add>
Customer Helpline	<Local market to add>
Fax	<Local market to add>

By completing the information below you confirm you have read, understood and distributed the contents of this Field Safety Notice accordingly.

2. Customer Details			
Healthcare Organisation / Facility Name*	<Fillable form field>		
Name of all Facilities/Hospitals covered by this response*	<Fillable form field>		
Facility / Hospital Address*	<Fillable form field>		
Telephone Number	<Fillable form field>	Email address	<Fillable form field>
Name of your supplier / wholesaler (if not Smith+Nephew)	<Fillable form field>		
Healthcare Organisation / Facility Stamp (if available)	<Fillable form field>		

3. Customer action undertaken on behalf of Healthcare Organisation / Facility Please complete/tick as appropriate.	
<input type="checkbox"/> Yes	I confirm receipt of the Field Safety Notice and that I read and understood its content.*
<input type="checkbox"/> Yes <input type="checkbox"/> No	Has your Healthcare Organisation / Facility distributed the product to other organisations? If you have answered yes, tick all that apply: *
	<input type="checkbox"/> I have identified customers that received or may have received this device.
	<input type="checkbox"/> I have informed the identified customers of this FSN.
<input type="checkbox"/> Yes	I have received confirmation of reply from all identified customers.
<input type="checkbox"/> Yes	I performed all actions requested by the FSN. *
Tick Appropriate Response:*	<input type="checkbox"/> Yes Neither I nor any of my customers has any affected devices in inventory.
	<input type="checkbox"/> Yes In our Organisation / Facility we have concerned devices that: <ul style="list-style-type: none"> - have been placed in quarantine and - returned as indicated in Section 4 below. Complete Section 4 with material, batch/serial, and quantity information related to devices to be returned.

4. Devices to be Returned		
Material Number	Batch or Serial Number	Quantity Quarantined and to be returned

Print Name*	<Fillable form field>		
Signature*	<Fillable form field>	Date*	<Fillable form field>

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 actions.