



[FDA Home](#)<sup>3</sup> [Medical Devices](#)<sup>4</sup> [Databases](#)<sup>5</sup>

## Class 2 Device Recall CRE Pro Wireguided 1820mm 240cm



[510\(k\)](#)<sup>7</sup> [DeNovo](#)<sup>8</sup> [Registration & Listing](#)<sup>9</sup> [Adverse Events](#)<sup>10</sup> [Recalls](#)<sup>11</sup> [PMA](#)<sup>12</sup> [HDE](#)<sup>13</sup> [Classification](#)<sup>14</sup> [Standards](#)<sup>15</sup>  
[CFR Title 21](#)<sup>16</sup> [Radiation-Emitting Products](#)<sup>17</sup> [X-Ray Assembler](#)<sup>18</sup> [Medsun Reports](#)<sup>19</sup> [CLIA](#)<sup>20</sup> [TPLC](#)<sup>21</sup>

[New Search](#)

[Back to Search Results](#)

### Class 2 Device Recall CRE Pro Wireguided 1820mm 240cm



|                                           |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
|-------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <b>Date Initiated by Firm</b>             | May 21, 2026                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              |
| <b>Date Posted</b>                        | July 01, 2026                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             |
| <b>Recall Status</b> <sup>1</sup>         | Open <sup>3</sup> , Classified                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |
| <b>Recall Number</b>                      | Z-2631-2026                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               |
| <b>Recall Event ID</b>                    | <a href="#">99067</a> <sup>23</sup>                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |
| <b>510(K)Number</b>                       | <a href="#">K112994</a> <sup>24</sup>                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     |
| <b>Product Classification</b>             | <a href="#">Stents, drains and dilators for the biliary ducts</a> <sup>25</sup> - <b>Product Code</b> <a href="#">FGE</a> <sup>26</sup>                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   |
| <b>Product</b>                            | CRE Pro Wireguided 18-20mm 240cm                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          |
| <b>Code Information</b>                   | UPN: M00558710; GTIN: 8714729797593; Lot No. 38031485, 38031486, 38031487, 38031488, 38031489, 38031490, 38031642, 38031643, 38031644, 38031645, 38043600, 38043601, 38043602, 38043603, 38043604, 38055580, 38055581, 38055582, 38099704, 38099705, 38099706, 38099707, 38099709, 38099710, 38099711, 38099712, 38099713, 38099714, 38099715, 38110376, 38110377, 38110378, 38110379, 38131547, 38131548, 38150880, 38150881, 38150882, 38150883, 38162872, 38162873, 38162874, 38162875, 38169202, 38169203, 3818825438188255, 38193520, 38241877, 38241878, 38258885, 38258886, 38258887, 38258888, 38258889, 38278151, 38285220, 38285221, 38306689, 38306692, 38338285, 38338288, 38338290, 38338292, 38338294, 38338296, 38338297, 38338299, 38338301, 38347330, 38462126, 38462127, 38462128, 38462129, 38479753, 38479754, 38553523, 38553524, 38553525, 38608098, 38608099, 38612875, 38612876, 38612877, 38612878, 38622479, 38622600, 38622601, 38622602, 38622603, 38642700, 38642701, 38642702, 38642722, 38650472, 38665039, 38665260, 38665261, 38665262, 38665263, 38678316, 38678317, 38678318, 38678319, 38678760, 38702327, 38702328, 38702329, 38702330, 38702331, 38702332, 38709525, 38709526, 38709527, 38709528, 38734736, 38742717, 38742718, 38745588, 38745589, 38745590, 38748246, 38748247, 38748248, 38808893, 38808894, 38808897, 38808899, 38809140, 38823455, 38823456, 38823457, 38874308, 38874309, 38874310, 38874311, 38877005, 38877006, 38886252, 38886253, 38886254, 38886255, 38946205, 38946207, 38946209, 38959786, 38964092, 38964093; Exp. November 23, 2028 March 29, 2029. |
| <b>FEI Number</b>                         | 3005099803                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |
| <b>Recalling Firm/ Manufacturer</b>       | Boston Scientific Corporation<br>100 Boston Scientific Way<br>Marlborough MA 01752-1234                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   |
| <b>For Additional Information Contact</b> | Renee Archie<br>508-683-4523                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              |
| <b>Manufacturer Reason for Recall</b>     | Potential sterile breach of the pouches in which devices are packaged.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    |
| <b>FDA Determined Cause</b> <sup>2</sup>  | Package design/selection                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  |
| <b>Action</b>                             | An Urgent Medical Device Removal - Immediate Action Required notification was mailed to consignees on 5/21/26. This notification informed consignees that Boston Scientific was conducting a removal of affected devices due to a potential sterile breach of product packaging (pouches). Consignees are instructed to immediately stop further use or distribution of product and quarantine them. Consignees are to                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    |

complete and return the provided Reply Verification Tracking Form and return the product to Boston Scientific per the provided instructions. Consignees are to share the recall notification with any healthcare provider that uses the affected devices and with organizations to which product was further distributed. Consignees with any questions are to contact their Boston Scientific representative.

**Quantity in Commerce** 81,543 units

**Distribution** Worldwide distribution - US Nationwide and the countries of Australia, Austria, Bahrain, Bangladesh, Belgium, Bulgaria, Canada, Chile, China, Costa Rica, Croatia, Czech Republic, Denmark, Dominican Republic, Ecuador, Estonia, Finland, France, Germany, Great Britain, Greece, Guatemala, Hong Kong, Hungary, India, Indonesia, Ireland, Israel, Italy, Japan, Jordan, Latvia, Lebanon, Luxembourg, Malaysia, Mexico, Mongolia, Netherlands, New Zealand, Norway, Pakistan, Paraguay, Peru, Philippines, Poland, Portugal, Romania, Russian Federation, Serbia, Singapore, Slovakia, Slovenia, South Africa, South Korea, Spain, Sweden, Switzerland, Taiwan, Tobago, Trinidad, Tunisia, Türkiye, United Arab Emirates, Uruguay.

**Total Product Life Cycle** [TPLC Device Report](#)<sup>27</sup>

<sup>1</sup> A record in this database is created when a firm initiates a correction or removal action. The record is updated if the FDA identifies a violation and classifies the action as a recall, and it is updated for a final time when the recall is terminated. Learn more about [medical device recalls](#)<sup>28</sup>.

<sup>2</sup> Per FDA policy, recall cause determinations are subject to modification up to the point of termination of the recall.

<sup>3</sup> The manufacturer has initiated the recall and not all products have been corrected or removed. This record will be updated as the status changes.

**510(K) Database** [510\(K\)s with Product Code = FGE](#)<sup>29</sup>

#### Links on this page:

1. <http://www.addthis.com/bookmark.php?u508=true&v=152&username=fdmain>
2. <http://www.addthis.com/bookmark.php>
3. <https://www.fda.gov/>
4. <https://www.fda.gov/Medical-Devices>
5. <https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/medical-device-databases>
6. </scripts/cdrh/devicesatfda/index.cfm>
7. </scripts/cdrh/cfdocs/cfPMN/pmn.cfm>
8. </scripts/cdrh/cfdocs/cfpmn/denovo.cfm>
9. </scripts/cdrh/cfdocs/cfRL/rl.cfm>
10. </scripts/cdrh/cfdocs/cfMAUDE/TextSearch.cfm>
11. </scripts/cdrh/cfdocs/cfRES/res.cfm>
12. </scripts/cdrh/cfdocs/cfPMA/pma.cfm>
13. </scripts/cdrh/cfdocs/cfHDE/hde.cfm>
14. </scripts/cdrh/cfdocs/cfPCD/classification.cfm>
15. </scripts/cdrh/cfdocs/cfStandards/search.cfm>
16. </scripts/cdrh/cfdocs/cfCFR/CFRSearch.cfm>
17. [/scripts/cdrh/cfdocs/cfPCD\\_RH/classification.cfm](/scripts/cdrh/cfdocs/cfPCD_RH/classification.cfm)
18. </scripts/cdrh/cfdocs/cfAssem/assembler.cfm>
19. </scripts/cdrh/cfdocs/Medsun/searchReportText.cfm>
20. </scripts/cdrh/cfdocs/cfClia/Search.cfm>
21. </scripts/cdrh/cfdocs/cfTPLC/tpic.cfm>
22. <https://www.fda.gov/safety/recalls-market-withdrawals-safety-alerts/enforcement-reports>
23. [/scripts/cdrh/cfdocs/cfRES/res.cfm?start\\_search=1&event\\_id=99067](/scripts/cdrh/cfdocs/cfRES/res.cfm?start_search=1&event_id=99067)
24. </scripts/cdrh/cfdocs/cfPMN/pmn.cfm?ID=K112994>
25. </scripts/cdrh/cfdocs/cfPCD/classification.cfm?ID=FGE>
26. </scripts/cdrh/cfdocs/cfPCD/classification.cfm?ID=FGE>
27. </scripts/cdrh/cfdocs/cfTPLC/tpic.cfm?id=FGE>
28. <https://www.fda.gov/medical-devices/medical-device-recalls/what-medical-device-recall>

29. /scripts/cdrh/cfdocs/cfPMN/pmn.cfm?  
start\_search=1&productcode=FGE&knumber=&applicant=Boston%20Scientific%20Corporation

Page Last Updated: 08/06/2026

Note: If you need help accessing information in different file formats, see [Instructions for Downloading Viewers and Players](#).

Language Assistance Available: [Español](#) | [繁體中文](#) | [Tiếng Việt](#) | [한국어](#) | [Tagalog](#) | [Русский](#) | [العربية](#) | [Kreyòl Ayisyen](#) | [Français](#) | [Polski](#) | [Português](#) | [Italiano](#) | [Deutsch](#) | [日本語](#) | [فارسی](#) | [English](#)

[Accessibility Contact](#) [FDA Careers](#) [FDA Basics](#) [FOIA](#) [No FEAR Act](#) [Nondiscrimination](#) [Website Policies](#) / [Privacy](#)



U.S. Food and Drug Administration  
10903 New Hampshire Avenue  
Silver Spring, MD 20993  
Ph. 1-888-INFO-FDA (1-888-463-6332)  
[Contact FDA](#)



[For Government For Press](#)

[Combination Products](#) [Advisory Committees](#) [Science & Research](#) [Regulatory Information](#) [Safety](#) [Emergency Preparedness](#)  
[International Programs](#) [News & Events](#) [Training and Continuing Education](#) [Inspections/Compliance](#) [State & Local Officials](#)  
[Consumers](#) [Industry Health Professionals](#) [FDA Archive](#) [Vulnerability Disclosure Policy](#)



U.S. Department of **Health & Human Services**

#### Links on this page:

1. <http://www.addthis.com/bookmark.php?u508=true&v=152&username=fdomain>
2. <http://www.addthis.com/bookmark.php>
3. <https://www.fda.gov/>
4. <https://www.fda.gov/Medical-Devices>
5. <https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/medical-device-databases>
6. </scripts/cdrh/devicesatfda/index.cfm>
7. </scripts/cdrh/cfdocs/cfPMN/pmn.cfm>
8. </scripts/cdrh/cfdocs/cfpmn/denovo.cfm>
9. </scripts/cdrh/cfdocs/cfRL/rl.cfm>
10. </scripts/cdrh/cfdocs/cfMAUDE/TextSearch.cfm>
11. </scripts/cdrh/cfdocs/cfRES/res.cfm>
12. </scripts/cdrh/cfdocs/cfPMA/pma.cfm>
13. </scripts/cdrh/cfdocs/cfHDE/hde.cfm>
14. </scripts/cdrh/cfdocs/cfPCD/classification.cfm>
15. </scripts/cdrh/cfdocs/cfStandards/search.cfm>
16. </scripts/cdrh/cfdocs/cfCFR/CFRSearch.cfm>
17. [/scripts/cdrh/cfdocs/cfPCD\\_RH/classification.cfm](/scripts/cdrh/cfdocs/cfPCD_RH/classification.cfm)
18. </scripts/cdrh/cfdocs/cfAssem/assembler.cfm>
19. </scripts/cdrh/cfdocs/Medsun/searchReportText.cfm>
20. </scripts/cdrh/cfdocs/cfClia/Search.cfm>
21. </scripts/cdrh/cfdocs/cfTPLC/tpic.cfm>
22. <https://www.fda.gov/safety/recalls-market-withdrawals-safety-alerts/enforcement-reports>
23. [/scripts/cdrh/cfdocs/cfRES/res.cfm?start\\_search=1&event\\_id=99067](/scripts/cdrh/cfdocs/cfRES/res.cfm?start_search=1&event_id=99067)
24. </scripts/cdrh/cfdocs/cfPMN/pmn.cfm?ID=K112994>
25. </scripts/cdrh/cfdocs/cfPCD/classification.cfm?ID=FGE>
26. </scripts/cdrh/cfdocs/cfPCD/classification.cfm?ID=FGE>
27. </scripts/cdrh/cfdocs/cfTPLC/tpic.cfm?id=FGE>
28. <https://www.fda.gov/medical-devices/medical-device-recalls/what-medical-device-recall>
29. [/scripts/cdrh/cfdocs/cfPMN/pmn.cfm?  
start\\_search=1&productcode=FGE&knumber=&applicant=Boston%20Scientific%20Corporation](/scripts/cdrh/cfdocs/cfPMN/pmn.cfm?start_search=1&productcode=FGE&knumber=&applicant=Boston%20Scientific%20Corporation)