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3 On Behalf of Brian Martini (Plaintiff)  
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11 **UNITED STATES DISTRICT COURT**  
12 **CENTRAL DISTRICT OF CALIFORNIA**

13 BRIAN MARTINI

Case No. 2:26-cv-00825

14 v.

15 BOSTON SCIENTIFIC  
16 CORPORATION; UNITED  
17 STATES FOOD AND DRUG  
18 ADMINISTRATION

**COMPLAINT – ACTION SEEKING  
NATIONWIDE RELIEF**

19 **COMPLAINT – ACTION SEEKING NATIONWIDE RELIEF**

20 Plaintiff, by and through undersigned counsel, brings this First Amended  
21 Complaint against Defendants Boston Scientific Corporation and the United States  
22 Food and Drug Administration and alleges as follows:

23 **I. PARTIES, JURISDICTION, AND VENUE**

24 1. Plaintiff Brian Martini (“Plaintiff”) is and was at all relevant times a  
25 resident of Florida. Plaintiff was implanted with a spinal cord stimulator (“SCS”)  
26 system designed, manufactured, and distributed by Defendant Boston Scientific  
27 Corporation.  
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1           2. Defendant Boston Scientific Corporation (“Boston Scientific”) is a  
2 corporation organized under the laws of the State of Delaware. Boston Scientific is  
3 registered and interacts with the Food and Drug Administration through its  
4 offices located at 25155 Rye Canyon Loop, Valencia, CA 91355. Boston Scientific  
5 conducts business nationwide and within this District, including the design,  
6 manufacture, regulatory submission, and distribution of Class III  
7 neuromodulation devices, such as its spinal cord stimulator systems marketed  
8 under the trade names Precision Montage MRI and other similar devices.

9           3. Defendant United States Food and Drug Administration (FDA) is an  
10 agency of the United States government responsible for regulating medical  
11 devices under the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. § 301 et seq.,  
12 and its implementing regulations. The FDA is named solely in its official capacity  
13 for purposes of claims brought under the Administrative Procedure Act (“APA”), 5  
14 U.S.C. §§ 701–706.

15           4. Venue is proper in this District pursuant to 28 U.S.C. § 1391(b)(1)–  
16 (2) because Defendant Boston Scientific resides in this District and a substantial  
17 part of the events or omissions giving rise to the claims occurred here. Venue is  
18 also appropriate with respect to the FDA under 28 U.S.C. § 1391(e)(1) because a  
19 substantial part of the events giving rise to the APA claims occurred in this  
20 District.

21           5. This Court has subject matter jurisdiction over Plaintiff’s claims  
22 against the FDA pursuant to 28 U.S.C. § 1331 and 5 U.S.C. § 702. This Court has  
23 supplemental jurisdiction over related state-law claims pursuant to 28 U.S.C. §  
24 1367 and diversity jurisdiction under 28 U.S.C. § 1332, as the amount in  
25 controversy exceeds \$75,000 and the parties are citizens of different states.

26           6. This Court has personal jurisdiction over Boston Scientific because it  
27 maintains significant offices and personnel in this District and regularly conducts  
28 and solicits business, and derives substantial revenue from the design,

1 manufacture, and sale of spinal cord stimulators and related services within this  
2 District.

3 **II. FACTUAL ALLEGATIONS REGARDING BOSTON SCIENTIFIC**  
4 **SCS DEVICES AND REGULATORY HISTORY**

5 **A. Overview of Spinal Cord Stimulation Devices and Their Intended Use**

6 7. SCS devices are Class III implantable neuromodulation systems  
7 designed to deliver electrical impulses to the spinal cord to mask or modulate  
8 chronic intractable pain. SCS systems typically consist of an implantable pulse  
9 generator (IPG), one or more electrical leads, and external patient controllers for  
10 adjusting therapeutic levels.

11 8. The underlying therapeutic premise of SCS devices is that electrical  
12 stimulation of the dorsal columns can “override” or “mask” the transmission of  
13 pain signals to the brain, thereby providing relief for chronic pain conditions that  
14 are otherwise resistant to conventional treatments.

15 9. SCS devices have long been associated with complex risks, including  
16 but not limited to device migration, lead breakage, battery failure, infection,  
17 stimulation-induced neurological deficits, exacerbation of pain, and autonomic  
18 dysfunction.

19 10. Due to these inherent risks, SCS devices are classified by the FDA as  
20 Class III medical devices, requiring Premarket Approval (“PMA”) or PMA  
21 supplement review for any design or functional changes affecting the safety and  
22 effectiveness of the device.

23 11. These devices are also associated with significant complications and  
24 poor clinical results, including inadequate effectiveness in providing the pain  
25 relief they promise. In September 2020, in response to a Public Citizen report, the  
26 FDA issued a letter to healthcare providers advising that, during the preceding  
27 four-year period, it had received 107,728 adverse event reports regarding spinal  
28

1 cord stimulators. The letter also disclosed 30,321 reports of unsatisfactory pain  
2 relief.

3 **B. Boston Scientific’s Device Portfolio and Regulatory Approval History**

4 12. The Federal Food, Drug & Cosmetic Act (“FDCA”) sets forth the  
5 requirements for Premarket Approval, which is necessary for the  
6 commercialization of a high-risk Class III device such as the device at issue in this  
7 case.

8 13. Among the aforesaid requirements is the requirement to provide the  
9 FDA with adequate clinical data to support a finding of sufficient safety and  
10 efficacy of the subject device.

11 14. The FDCA’s implementing regulations require that an applicant for  
12 premarket approval of a device submit, with respect to the device proposed to be  
13 marketed:

14 results of the clinical investigations involving human subjects with the  
15 device including clinical protocols, number of investigators and subjects per  
16 investigator, subject selection and exclusion criteria, study population,  
17 study period, safety and effectiveness data, adverse reactions and  
18 complications [...]

19 15. Boston Scientific’s spinal cord stimulator product line originates from  
20 and is, for all intents and purposes, predicated on PMA P030017, initially  
21 approved by the FDA in 2004 for its Precision Spinal Cord Stimulator System.

22 16. PMA P030017 was awarded despite the applicant’s failure to supply  
23 the data required by 21 CFR §814.20.

24 17. Instead, the referenced PMA was granted following submission of  
25 clinical data from “available peer reviewed published literature for similar  
26 implantable spinal cord stimulation (SCS) systems.”

27 18. Boston Scientific did not submit and the FDA did not consider  
28 clinical data or clinical evidence for the Precision system, or for any subsequent

1 BSC system that used the Precision as a predicate product for marketing  
2 purposes.

3 19. Thus, the defining parameter for the grant of premarket approval of a  
4 medical device has never been met with respect to the device at issue in this case,  
5 or any other Boston Scientific SCS system for that matter.

6 20. Since the original approval of P030017, Boston Scientific has  
7 introduced numerous subsequent models and upgrades under PMA supplements,  
8 including the Precision Plus, Precision Spectra, Spectra WaveWriter, WaveWriter  
9 Alpha, and Precision Montage MRI systems.

10 21. Since the original approval of P030017, Boston Scientific has filed  
11 362 supplements to P030017.

12 22. Boston Scientific's newer generations of devices incorporated  
13 significant modifications, including multiwaveform stimulation (simultaneous  
14 tonic, burst, and sub-perception modes), posture-adaptive programming,  
15 expanded electrode arrays, Bluetooth-enabled programming, and major revisions  
16 to battery architecture and lead designs.

17 23. None of these systems are supported by premarket clinical data  
18 supporting safety or effectiveness.

19 24. Boston Scientific aggressively marketed its Precision Montage MRI  
20 system and other upgraded models as offering superior pain relief through  
21 innovative stimulation patterns, despite the absence of independent premarket  
22 clinical testing validating the long-term safety and effectiveness of these  
23 substantial modifications.

24 **C. Material Changes to Device Architecture and Functionality**

25 25. Over time, Boston Scientific introduced substantial modifications to  
26 the originally approved Precision SCS system, including:

- 27 • The addition of simultaneous multiwaveform stimulation,  
28 including tonic, burst, and sub-perception programming;

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- The redesign of the implantable pulse generator battery system and the addition of Bluetooth-enabled wireless communication capabilities;
- The integration of posture-adaptive stimulation algorithms;
- The expansion of lead configurations and multi-source current delivery systems.

26. These substantial modifications materially altered the device's safety and effectiveness profile compared to the originally approved Precision system, triggering regulatory obligations that Boston Scientific failed to fulfill.

27. Under 21 C.F.R. § 814.39(a), such significant changes require submission of a new PMA or substantial clinical data demonstrating continued safety and effectiveness. Boston Scientific failed to pursue a new PMA review for these cumulative design changes and failed to provide substantial clinical data demonstrating continued safety and effectiveness, and instead improperly utilized the PMA supplement pathway.

28. Exemplifying the significant changes Boston Scientific has improperly made to its SCS systems over the years, it currently advertises and markets three distinct systems on its website – the Precision Montage, Spectra WaveWriter, and the WaveWriter Alpha systems. Boston Scientific advertises and markets these systems as distinct from each other. Despite advertising three distinct systems, it has only ever received one PMA approval.

29. In a May 19, 2016 press release, Boston Scientific described the Precision Montage MRI system as its “new SCS system.”

30. The Precision Montage MRI implanted in Plaintiff is technologically unrecognizable from the Precision system that was originally approved by the FDA in 2004.

1 31. Throughout its history in the neuromodulation space, BSC has  
2 conflated the PMA approval process with the less onerous and demanding 510k  
3 clearance process.

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5 **D. Regulatory Manipulation and Abuse of the PMA Supplement Process**

6 32. Boston Scientific submitted successive PMA supplements treating  
7 major modifications as discrete “minor” changes to avoid the heightened scrutiny,  
8 public transparency, and rigorous independent clinical evaluation required for  
9 new PMA applications.

10 33. This regulatory strategy deprived physicians, patients, and the FDA  
11 of complete information necessary to evaluate the true risks associated with the  
12 modified devices, particularly in the areas of neurological safety, device longevity  
13 device effectiveness, stimulation safety, and autonomic complications.

14 34. As a direct consequence of these omissions and regulatory  
15 manipulations, the Precision Montage MRI and other successor systems entered  
16 the market and were widely implanted without sufficient scientific validation of  
17 their safety and effectiveness.

18 **E. Post-Market Failures, Adverse Events, and Concealment of Risks**

19 35. Publicly available MAUDE (Manufacturer and User Facility Device  
20 Experience) database entries, peer-reviewed studies, and post-market surveillance  
21 data demonstrate that Boston Scientific’s SCS systems are associated with serious  
22 complications, including:

- 23 • Unsatisfactory pain relief;
- 24 • Device migration and loss of therapeutic coverage;
- 25 • Lead fractures requiring surgical revision;
- 26 • Battery depletion and communication failures;
- 27 • Stimulation-induced autonomic dysfunction, including urinary  
28 incontinence and orthostatic hypotension;

- Persistent ineffective pain relief despite extensive reprogramming.

36. Peer-reviewed literature has increasingly associated SCS therapy, particularly multiwaveform stimulation platforms like the Precision Montage MRI, with unexpected autonomic side effects. *See, e.g., Steven Smeijers et al., Spinal Cord Stimulation and Urinary Dysfunction, 23 Pain Med. 1204, 1204–1216 (2022).*

37. A 2023 Cochrane Review led by University of Sydney researchers found that, based on all randomized controlled trials and cross-over trials comparing SCS with placebo or no treatment for low back pain, current evidence suggests that SCS probably does not have sustained clinical benefits that would outweigh the costs and risks of this surgical intervention. *See Traeger AC, Gilbert SE, Harris IA, Maher CG. Spinal cord stimulation for low back pain. Cochrane Database of Systematic Reviews 2023, Issue 3. Art. No.: CD014789. DOI: 10.1002/14651858.CD014789.pub2. Accessed 08 January 2026.*

38. Despite knowledge of these adverse outcomes, Boston Scientific failed to timely update device labeling, issue field safety notices, or seek revised PMA approvals as required under federal regulations.

39. Of note, Boston Scientific has never conducted randomized, double-blind, placebo-controlled studies on any of its SCS devices.

40. Plaintiff’s injuries occurred as a direct and foreseeable result of Defendant Boston Scientific’s conduct and the FDA’s arbitrary and capricious regulatory approvals as set forth herein.

### **III. REGULATORY FRAMEWORK AND DUTIES**

41. Spinal cord stimulator (SCS) systems are regulated as Class III medical devices under the Federal Food, Drug, and Cosmetic Act (“FDCA”), 21 U.S.C. § 301 et seq., and the Medical Device Amendments of 1976 (“MDA”), 21 U.S.C. § 360c et seq.

1 42. Class III devices are those that present the highest risk to patients  
2 and are subject to the most rigorous form of regulatory oversight, including the  
3 requirement to obtain Premarket Approval from the FDA prior to marketing. *See*  
4 21 U.S.C. § 360e.

5 43. To obtain PMA, a manufacturer must submit detailed information  
6 demonstrating the safety and effectiveness of the device, including clinical trial  
7 data, descriptions of manufacturing methods, proposed labeling, and a risk-  
8 benefit analysis. *See* 21 C.F.R. § 814.20.

9 44. Once a PMA is granted, any proposed changes to the device's design,  
10 labeling, intended use, or manufacturing process must be submitted to the FDA as  
11 a PMA supplement. *See* 21 C.F.R. § 814.39(a).

12 45. The MDA distinguishes between different types of PMA supplements  
13 based on the nature and significance of the change. A “panel-track supplement” is  
14 required for changes that affect the safety or effectiveness of the device, such as  
15 new indications for use, major design modifications, or significant changes in  
16 component materials. *See* 21 C.F.R. § 814.39(c).

17 46. For any change that could affect safety or effectiveness, the FDA must  
18 receive and approve the PMA supplement before the manufacturer implements  
19 the change. *Id.* The burden of proof remains with the manufacturer to  
20 demonstrate that the modified device continues to be safe and effective.

21 47. Manufacturers are also required to comply with post-market  
22 surveillance and reporting obligations, including:

- 23 • Timely submission of adverse event reports under 21 C.F.R. §  
24 803.50;
- 25 • Maintenance of complaint files under 21 C.F.R. § 820.198;
- 26 • Evaluation of nonconforming products under 21 C.F.R. § 820.90;
- 27 • Implementation of corrective and preventive actions under 21  
28 C.F.R. § 820.100.

1 48. These regulatory obligations are non-discretionary and enforceable  
2 under both federal and state law. A manufacturer’s failure to comply with these  
3 requirements renders its device adulterated or misbranded under 21 U.S.C. §§ 351  
4 and 352.

5 49. Additionally, under the APA, 5 U.S.C. §§ 701–706, FDA actions that  
6 are arbitrary, capricious, an abuse of discretion, or otherwise not in accordance  
7 with law are subject to judicial review.

8 50. The FDA’s passive acceptance or failure to meaningfully review  
9 Boston Scientific’s PMA supplements—particularly where those supplements  
10 concealed the scope and safety implications of the changes—constitutes agency  
11 action unlawfully withheld and final agency action subject to review under 5  
12 U.S.C. §§ 706(1), 706(2)(A)–(D).

13 51. Plaintiff does not seek to enforce the FDCA directly. Rather, Plaintiff  
14 asserts state-law tort claims based on duties that parallel and incorporate federal  
15 requirements, as recognized in *Medtronic, Inc. v. Lohr*, 518 U.S. 470 (1996), and  
16 as preserved under 21 U.S.C. § 360k(a).

17 52. These state-law claims are not preempted by the MDA because they  
18 are premised on conduct that violates both federal law and equivalent duties  
19 imposed by Florida and California law. Plaintiff also seeks judicial review under  
20 the APA for final agency action by the FDA that facilitated or failed to correct the  
21 wrongful acts of Defendant Boston Scientific.

22 **IV. ALLEGATIONS AGAINST THE FDA UNDER THE**  
23 **ADMINISTRATIVE PROCEDURE ACT**

24 53. Plaintiff realleges and incorporates by reference all preceding  
25 paragraphs of this Complaint as though fully set forth herein.

26 54. Defendant United States Food and Drug Administration is an agency  
27 of the United States government charged with ensuring that medical devices  
28 marketed in the United States are safe and effective for their intended use,

1 pursuant to the FDCA, 21 U.S.C. § 301 *et seq.*, and the MDA, 21 U.S.C. § 360c *et*  
2 *seq.*

3 55. Under the APA, 5 U.S.C. §§ 701–706, federal courts are authorized to  
4 review final agency actions, including agency actions that are arbitrary,  
5 capricious, an abuse of discretion, or otherwise not in accordance with law. See 5  
6 U.S.C. §§ 706(1), 706(2)(A)–(D); *Loper Bright Enterprises v. Raimondo*, 603 U.S.  
7 369 (2024).

8 56. The FDA’s passive acceptance and approval of Boston Scientific’s  
9 original PMA P030017 and numerous PMA supplements submitted by Boston  
10 Scientific Corporation for spinal cord stimulator devices under PMA P030017,  
11 including the Precision Montage MRI system implanted in Plaintiff, constituted  
12 final agency action within the meaning of the APA.

13 57. In 2004, the FDA failed to require Boston Scientific to submit the  
14 clinical data required by 21 CFR §814.20 in support of PMA P030017.

15 58. Instead, the FDA granted PMA P030017 despite Boston Scientific’s  
16 failure to provide the required data, instead granting approval expressly based  
17 only on literature, not clinical data, for similar implantable spinal cord  
18 stimulation systems.

19 59. The FDA also failed to require Boston Scientific to submit new  
20 Premarket Approval applications for substantial modifications to the original  
21 Precision SCS System, including:

- 22 • The addition of simultaneous multiwaveform stimulation (tonic,  
23 burst, sub-perception);
- 24 • The redesign of the IPG battery architecture and the addition of  
25 Bluetooth-enabled wireless communication features;
- 26 • The integration of posture-adaptive stimulation algorithms;
- 27 • The expansion of lead configurations and multi-source current  
28 delivery systems.

1           60. By approving substantial cumulative changes via successive PMA  
2 supplements without requiring full panel-track review or independent clinical  
3 validation, the FDA unlawfully allowed Boston Scientific to materially alter the  
4 design, intended use, and safety profile of its spinal cord stimulator systems  
5 outside the bounds of statutory and regulatory requirements. *See* 21 U.S.C. §  
6 360e(d); 21 C.F.R. § 814.39(a).

7           61. Essentially, the FDA allowed Boston Scientific to bring new Class III  
8 SCS products, including the original Precision SCS and subsequent SCS products,  
9 to market based on predicate products. This process much more closely resembles  
10 the FDA's 510(k) premarket submission process than the Premarket Approval  
11 process. A critical difference between these two routes, however, is that the PMA  
12 process gives BSC the ability to raise federal preemption as a defense to Plaintiff's  
13 claims.

14           62. The FDA's acceptance and approval of Boston Scientific's original  
15 PMA P030017 and subsequent PMA supplements submitted by Boston Scientific  
16 constitutes "final agency action"<sup>1</sup> under 5 U.S.C. § 551(13) and 5 U.S.C. § 704. A  
17 "final agency action" is one that (1) marks the consummation of the agency's  
18 decision-making process and (2) determines rights or obligations or from which  
19 legal consequences flow. *See Bennett v. Spear*, 520 U.S. 154, 177–78 (1997). The  
20 FDA's approval of PMA P030017 and subsequent PMA supplements was the final  
21 step in the regulatory process, authorizing the commercial marketing of spinal  
22 cord stimulators that were not supported by clinical evidence of safety or  
23 effectiveness, and subsequently, of materially modified spinal cord stimulator  
24 systems that were equally unsupported by clinical evidence of safety or

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26 <sup>1</sup> Under the Administrative Procedure Act, "agency action" includes "the whole or a part  
27 of an agency rule, order, license, sanction, relief, or the equivalent or denial thereof, or  
28 failure to act," and final agency action is judicially reviewable when it consummates the  
decision-making process and determines rights or obligations or produces legal  
consequences. *See* 5 U.S.C. § 551(13); 5 U.S.C. § 704; *Bennett v. Spear*, 520 U.S. 154,  
177–78 (1997).

1 effectiveness. This approval carried immediate and direct legal consequences,  
2 allowing Boston Scientific to market and distribute altered devices nationwide  
3 under federal premarket authorization, behind the legal shield of federal  
4 preemption. Plaintiff's Administrative Procedure Act claims challenge these final  
5 agency actions, not discretionary enforcement decisions or ongoing regulatory  
6 processes, and thus fall squarely within the scope of judicial review authorized by  
7 5 U.S.C. §§ 702 and 706.

8 63. The FDA's actions and omissions materially contributed to the  
9 injuries suffered by Plaintiff by enabling the marketing and widespread  
10 implantation of devices whose risks had not been fully evaluated or disclosed,  
11 depriving physicians and patients of critical safety information.

12 64. The FDA's actions and omissions materially contributed to Plaintiff's  
13 legal injury, in that these actions and omissions allow Boston Scientific to invoke  
14 federal preemption to argue that Plaintiff should be barred from seeking redress  
15 for physical injuries caused by its SCS system.

16 65. But for the FDA's actions and omissions, Boston Scientific would not  
17 have received PMA approval for the SCS system implanted in Plaintiff, and  
18 therefore, would not be able to invoke federal preemption to shield itself from  
19 liability for the physical injuries that the SCS system caused Plaintiff.

20 **Evidence of Agency Capture: ANA Reclassification Petition and FDA**  
21 **Override**

22 66. In 2001, Advanced Neuromodulation Systems (ANS) submitted a  
23 petition to the FDA requesting that implantable spinal cord stimulators be  
24 reclassified from Class III to Class II devices. *See* FDA Docket No. 02P-0321.

25 67. An FDA advisory panel, after reviewing the available scientific  
26 evidence, recommended granting the reclassification request, concluding that SCS  
27 devices did not warrant Class III treatment based on their risk profiles and clinical  
28 experience.

1           68. Despite the advisory panel’s recommendation, FDA headquarters  
2 overruled the panel. They unilaterally decided to maintain Class III classification  
3 for SCS devices, yet without imposing meaningful PMA enforcement obligations  
4 or new clinical evidence requirements thereafter.

5           69. This historical regulatory decision is evidence of agency capture and  
6 arbitrary administrative conduct. The FDA simultaneously acknowledged that  
7 SCS devices did not merit full Class III regulatory burdens, yet failed to require  
8 proper PMA oversight for subsequent generations of increasingly complex and  
9 modified SCS systems.

10           70. The FDA’s passive and deferential treatment of Boston Scientific’s  
11 original PMA Application and subsequent PMA supplements following the  
12 override of its advisory panel further illustrates systemic regulatory failure and  
13 arbitrary decision-making in violation of the APA.

14           71. Plaintiff was injured as a direct and foreseeable result of the FDA’s  
15 arbitrary and unlawful regulatory actions. Had the FDA properly enforced PMA  
16 standards and statutory requirements, the Precision Montage MRI device  
17 implanted in Plaintiff would not have entered the market at all, and certainly not  
18 in the materially altered form that was ultimately implanted in Plaintiff. Further,  
19 Boston Scientific would not be able to invoke the defense of federal preemption to  
20 shield itself from liability for the physical injuries that the Precision Montage MRI  
21 caused Plaintiff.

22           72. Plaintiff seeks judicial review of these final agency actions under the  
23 APA, including declaratory and injunctive relief as necessary to vindicate  
24 statutory rights, remedy the injury caused by the agency’s conduct, and prevent  
25 ongoing harm to Plaintiff and the public.

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1 **V. PLAINTIFF-SPECIFIC ALLEGATIONS**

2 73. Plaintiff Brian Martini is a resident of Destin, Florida. On November  
3 7, 2018, Plaintiff underwent implantation of a Boston Scientific spinal cord  
4 stimulator system.

5 74. The implanted system included a Precision Montage MRI  
6 implantable pulse generator, along with Boston Scientific leads and anchoring  
7 components.

8 75. Plaintiff's spinal cord stimulator was implanted for the treatment of  
9 chronic pain.

10 76. Prior to deciding to have the Precision Montage MRI system  
11 implanted, Plaintiff met multiple times with Boston Scientific sales  
12 representatives, including Wayne (last name unknown).

13 77. Throughout the time period starting with his first contact with a  
14 Boston Scientific representative and ending with the removal of the SCS system,  
15 Plaintiff had contact with four Boston Scientific representatives.

16 78. Before being permanently implanted with the Precision Montage  
17 MRI system, Plaintiff underwent a temporary spinal cord stimulator trial.

18 79. Plaintiff was covered by Medicare at the time he was implanted with  
19 the SCS system, and the cost of the SCS system was paid by Medicare.

20 80. For Medicare to pay for the cost of the SCS system, Plaintiff and his  
21 physician had to report a certain level of pain relief from the trial stimulator.  
22 Thus, statements made to Plaintiff regarding the perceived effectiveness of the  
23 trial, and any purported predictions about the effectiveness of the permanent SCS  
24 system were critical to the insurance coverage of the very expensive permanent  
25 implant.

26 81. Prior to the SCS trial, Plaintiff was advised by Boston Scientific sales  
27 representatives that the permanent SCS device would provide reliable, long-term  
28 pain relief and that stimulation was safe and localized. No warnings were given

1 regarding the risk of shocking, the possibility that the permanent device differed  
2 materially from the external trial unit, or the likelihood that the SCS system would  
3 not provide meaningful, long-term pain relief. In fact, the Boston Scientific  
4 representative represented to Plaintiff that the permanent SCS system would  
5 provide equivalent pain relief to the temporary SCS trial, and would provide  
6 consistent long-term pain relief.

7 82. During the SCS trial, Plaintiff communicated with a Boston Scientific  
8 representative regarding the efficacy of the trial stimulator unit.

9 83. At the end of the SCS trial, Plaintiff reported greater than eighty  
10 percent pain relief.

11 84. Plaintiff and his providers were not informed that the original PMA  
12 that led to the Precision Montage MRI system being on the market (PMA  
13 P030017) was not supported by clinical trial data or that it was approved based on  
14 literature relating to other spinal cord stimulator devices.

15 85. Plaintiff and his providers were also not informed that the Precision  
16 Montage MRI system, including its multiwaveform stimulation capabilities and  
17 revised battery architecture, had not undergone independent clinical validation  
18 via a new PMA application, and instead reached the market through piecemeal  
19 PMA supplement filings.

20 86. Plaintiff relied on Boston Scientific's representations, through both  
21 direct promotional materials and the direct spoken representations by the Boston  
22 Scientific sales representatives in deciding to proceed with the initial implantation  
23 of the Precision Montage MRI system.

24 87. Specifically, Plaintiff relied on Boston Scientific's representations,  
25 among others, that the Precision Montage MRI system was uniquely suited to  
26 provide him with long-term pain relief and was safe for long-term implantation,  
27 that the system was supported by clinical data, and that the permanent SCS  
28 system would provide equivalent pain relief to the temporary SCS trial.

1 88. The representations relied upon by Plaintiff are still being made by  
2 Boston Scientific about the Precision Montage MRI to this day.

3 89. Plaintiff's interactions with Boston Scientific's representatives were  
4 so substantial that Plaintiff alleges that they directly sold him the Precision  
5 Montage MRI system.

6 90. A Boston Scientific representative was present at the surgery facility  
7 on the day of Plaintiff's permanent implant surgery, and programmed the device  
8 in the PACU immediately after the implant surgery.

9 91. On approximately ten occasions, Plaintiff was required to undergo  
10 reprogramming of the SCS device by Boston Scientific representatives due to  
11 unsatisfactory pain relief and severe pain, electric shocking sensations, and left  
12 lower extremity pain.

13 92. After some of these reprogramming sessions, Plaintiff did receive  
14 some temporary relief of his underlying pain and complications.

15 93. On the occasions that he had the SCS device reprogrammed, the  
16 Boston Scientific sales representatives represented to Plaintiff that  
17 reprogramming of the SCS system after it was implanted was necessary to ensure  
18 that he received optimal pain relief and to avoid the side effects he was  
19 experiencing, and that if he was not receiving adequate pain relief or experiencing  
20 these side effects, he just needed to have the system reprogrammed. Therefore,  
21 Plaintiff did not have any reason to know that he should not expect the Precision  
22 Montage MRI device to work properly and provide the promised results on the  
23 dates that he had the device reprogrammed.

24 94. On June 4, 2020, Plaintiff underwent a surgery to move the SCS  
25 system battery due to site pain.

26 95. On multiple occasions while the SCS system was implanted in  
27 Plaintiff, he felt that the battery was warming and becoming uncomfortably hot  
28 while it was charging.

1 96. On February 1, 2022, Plaintiff underwent a surgery to have the SCS  
2 device removed from his body permanently.

3 97. The permanent Precision Montage MRI system did not provide  
4 equivalent pain relief to Plaintiff's temporary SCS trial.

5 98. During the time in which the SCS system was implanted in Plaintiff,  
6 Boston Scientific representatives, believed to be unlicensed in the State of Florida,  
7 or elsewhere for that matter, actively participated in programming and waveform  
8 selection. These actions involved real-time interpretation of patient responses and  
9 materially influenced the configuration and function of the implanted system.  
10 These actions were essentially medical treatment and had a significant impact on  
11 the way the SCS system affected Plaintiff's body.

12 99. Plaintiff was advised that Boston Scientific representatives were the  
13 only individuals who could or would program or reprogram his SCS device.

14 100. During the time in which the SCS system was implanted in Plaintiff,  
15 Boston Scientific representatives provided medical advice to him about the SCS  
16 system.

17 101. After the SCS system was removed, Plaintiff continued to experience  
18 pain, including left lower extremity pain. He experiences this pain to this day, due  
19 to nerve damage caused by the SCS system.

20 102. Plaintiff relied on representations made and advice given by Boston  
21 Scientific representatives after he was implanted with the SCS system in electing  
22 to keep the system implanted in his body. If not for these representations and the  
23 purported medical advice provided by the Boston Scientific representatives,  
24 Plaintiff would have elected to have the system removed earlier and would have  
25 avoided much of the pain and suffering that he ultimately endured. Specifically, if  
26 the representatives had not convinced him to keep the faulty device in his body,  
27 he would have had it removed immediately and would likely have avoided the  
28 nerve damage he ultimately suffered.

1           103. Plaintiff did not learn of the likely connection between his injuries  
2 and the defective nature of Boston Scientific’s Precision Montage MRI system  
3 until he reviewed publicly available FDA documents, adverse event disclosures,  
4 and litigation-related findings that contradicted prior assurances of safety,  
5 efficacy, and regulatory compliance. Further, he did not and could not have  
6 known that any of Boston Scientific’s representations were false until he  
7 ultimately had the SCS system removed on February 1, 2022.

8           104. Boston Scientific’s tortious conduct continued through the date on  
9 which Plaintiff ultimately had the Boston Scientific SCS system removed from his  
10 body, as Boston Scientific sales representatives continued to make  
11 misrepresentations to him, continued to reprogram the SCS system, and gave  
12 medical advice to him about the SCS system.

13           105. Plaintiff’s injuries, including physical pain, emotional distress,  
14 surgical trauma, loss of enjoyment of life, and the permanent implantation of  
15 defective hardware, were directly and proximately caused by the acts and  
16 omissions of Boston Scientific, as well as the FDA’s unlawful and arbitrary failure  
17 to require a new PMA for the substantially modified Precision Montage MRI  
18 device.

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20 **VI. DEFENDANTS’ MISREPRESENTATIONS, OMISSIONS, AND**  
21 **REGULATORY VIOLATIONS**

22 **A. Failure to Disclose Material Risks and Regulatory Evasion**

23           106. Plaintiff realleges and incorporates by reference all preceding  
24 paragraphs of this Complaint as though fully set forth herein.

25           107. At all relevant times, Defendant Boston Scientific Corporation  
26 engaged in a course of conduct designed to conceal material risks associated with  
27 its spinal cord stimulator systems, misrepresent the safety and efficacy of its  
28 devices, and improperly utilize the PMA supplement process to introduce

1 significant, unvalidated design changes without triggering mandatory premarket  
2 review.

3 108. Boston Scientific represented to Plaintiff, his healthcare providers,  
4 and the medical community that its Precision Montage MRI system and related  
5 devices were safe, effective, supported by clinical data, and appropriately  
6 approved for long-term implantation.

7 109. These representations were false, misleading, and incomplete. Boston  
8 Scientific knew, or should have known through post-market surveillance and  
9 regulatory obligations, that the Precision Montage MRI system:

- 10 • Was not supported by clinical data;
- 11 • Posed an increased risk of device migration, stimulation failure, and  
12 neurological injury;
- 13 • Was marketed with stimulation modalities not adequately tested in  
14 clinical trials;
- 15 • Carried a known risk of autonomic dysfunction, including  
16 incontinence, hypotension, and cardiac arrhythmia;
- 17 • Had materially different performance characteristics from the  
18 external trial device, upon which ultimate treatment predictions and  
19 decisions were made.

20 110. Boston Scientific actively concealed this information by:

- 21 • Failing to conduct clinical trials and submit meaningful clinical  
22 data to the FDA as required by 21 U.S.C. § 360e(d) and 21 CFR §  
23 814.20.
- 24 • Failing to report adverse events under 21 C.F.R. § 803.50;
- 25 • Withholding labeling updates and field safety communications  
26 under 21 C.F.R. § 814.39(d);  
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- Submitting fragmented PMA supplements to avoid full panel-track review required by 21 C.F.R. § 814.39(a).

111. These actions violated non-discretionary regulatory obligations and rendered the device unsafe for its intended use and deprived physicians and patients of information required for informed medical decision-making.

**B. Violations of Current Good Manufacturing Practices (cGMPs)**

112. In addition to the above, Boston Scientific violated multiple cGMP requirements codified at 21 C.F.R. Part 820 — including those governing design control, process validation, complaint handling, and corrective and preventive action (CAPA).

113. Specifically, Boston Scientific:

- Failed to maintain adequate design validation under 21 C.F.R. § 820.30(g), particularly in light of the software and waveform changes introduced with the Precision Montage MRI system;
- Failed to validate and monitor manufacturing processes as required under 21 C.F.R. § 820.75, leading to inconsistencies in lead bonding and IPG housing integrity;
- Failed to investigate and correct known device performance issues through its CAPA system, in violation of 21 C.F.R. § 820.100;
- Failed to evaluate post-market complaints systematically and incorporate them into product redesign and labeling revisions, in violation of 21 C.F.R. § 820.198.

114. These cGMP violations are not discretionary; they are binding legal obligations imposed by the FDA to ensure the safety and effectiveness of devices. They establish minimum standards for medical device manufacturers and are incorporated by reference into Florida tort law as parallel duties.

115. Boston Scientific's violation of these cGMP requirements caused Plaintiffs' injuries described herein. It was reasonably foreseeable that these

1 violations would cause injury to Plaintiff and others. Specifically, these violations  
2 resulted in an unsafe product being marketed to Plaintiff and deprived Plaintiff  
3 and her physicians of important information about the safety and efficacy of the  
4 device at issue.

5 116. The Ninth Circuit has expressly recognized that manufacturers of  
6 FDA-regulated medical devices may be held liable under state law where they  
7 violate non-discretionary FDA regulations, including those arising under cGMPs.  
8 *See Stengel v. Medtronic Inc.*, 704 F.3d 1224, 1233–34 (9th Cir. 2013) (*en banc*).

9 117. Boston Scientific’s repeated and willful violations of cGMP  
10 requirements caused or substantially contributed to the defects and injuries at  
11 issue in this case, and support Plaintiff’s claims under Florida law.

12 118. These claims are not preempted under *Riegel* or *Buckman* because  
13 they are premised on state-law duties that genuinely parallel federal requirements  
14 and do not exist solely by virtue of the FDCA.

## 15 **VII. CAUSES OF ACTION**

### 16 **COUNT I – MANUFACTURING DEFECT**

#### 17 *Against Defendant Boston Scientific*

18 (Florida Common Law; 21 U.S.C. § 360k(a); 21 C.F.R. §§ 820.30, 820.70, 820.75,  
19 820.100)

20 119. Plaintiff realleges and incorporates by reference all preceding  
21 paragraphs of this Complaint as though fully set forth herein.

22 120. At all relevant times, Defendant Boston Scientific Corporation was  
23 engaged in the design, manufacture, labeling, marketing, and distribution of  
24 spinal cord stimulator systems throughout the United States, including the  
25 Precision Montage MRI system implanted in Plaintiff.

26 121. Under Florida law, a product is unreasonably dangerous because of a  
27 manufacturing defect if it was not manufactured according to its intended design  
28 and fails to perform as safely as an ordinary consumer would expect.

1 122. The spinal cord stimulator system implanted in Plaintiff was not  
2 reasonably safe as manufactured. It deviated materially from its intended design  
3 specifications and from applicable federal requirements governing Class III  
4 medical devices.

5 123. Defendant Boston Scientific violated non-discretionary federal  
6 manufacturing and quality system regulations, including but not limited to:

- 7 • 21 C.F.R. § 820.30: failure to implement adequate design controls;
- 8 • 21 C.F.R. § 820.70: failure to establish process controls ensuring  
9 conformity to design specifications;
- 10 • 21 C.F.R. § 820.75: failure to validate manufacturing processes  
11 capable of consistently producing conforming devices;
- 12 • 21 C.F.R. § 820.100: failure to implement corrective and  
13 preventive action in response to known device defects.

14 124. The Precision Montage MRI system implanted in Plaintiff was  
15 manufactured with latent defects affecting performance, stability, and safety,  
16 including but not limited to:

- 17 • Inadequate lead anchoring, resulting in migration, therapeutic  
18 failure, and unwanted shocking in patients;
- 19 • Battery instability contributing to irregular stimulation output and  
20 early depletion, including need for battery replacement;
- 21 • Defects impacting the physical stability of the battery in the bodies  
22 of patients;
- 23 • Faulty firmware or programming inconsistencies affecting  
24 waveform delivery, and causing complications in patients,  
25 including shocking and unsatisfactory pain relief.

26 125. These defects were not detectable by Plaintiff or her physicians  
27 through reasonable inspection prior to implantation and were not reasonably  
28 foreseeable based on the information available to Plaintiff or her treating  
physicians at the time.



1           131. Boston Scientific had a duty to warn Plaintiff, her healthcare  
2 providers, and the medical community of the material risks associated with its  
3 spinal cord stimulator systems, including but not limited to:

- 4           • The risk of lead migration and device failure requiring surgical  
5 revision;
- 6           • The risk of autonomic dysfunction, including arrhythmias, urinary  
7 incontinence, and hypotension, and neurological injury;
- 8           • The risk that the trial device would not reliably predict the  
9 performance of the permanently implanted device;
- 10          • The risk of device-related pain exacerbation, shocking  
11 complications, nerve damage, and loss of therapeutic efficacy over  
12 time.

13           132. Boston Scientific breached its duty to warn by:

- 14           • Failing to update product labeling and Instructions for Use (IFU)  
15 to reflect emerging adverse event trends in the nearly twenty years  
16 since PMA P030017 was initially approved;
- 17           • Failing to disseminate “Dear Doctor” letters or field advisories  
18 warning of known failure modes that arose since PMA P030017  
19 was approved;
- 20           • Failing to adequately train sales representatives and clinicians  
21 regarding the safety and efficacy limitations of multiwaveform  
22 stimulation, and the Precision Montage MRI system generally;
- 23           • Actively promoting the Precision Montage MRI system as  
24 superior, effective, and safe without disclosing material  
25 limitations and risks;
- 26           • Actively promoting the permanent Precision Montage MRI as  
27 superior or equivalent in efficacy to temporary trial SCS.

28           133. These failures were compounded by violations of federal law,  
including:

- 1 • 21 C.F.R. § 803.50: failure to timely report adverse events;
- 2 • 21 C.F.R. § 814.39: failure to submit PMA supplements for
- 3 significant changes;
- 4 • 21 C.F.R. § 820.198: failure to investigate and address post-
- 5 market complaints.

6 134. Plaintiff and her physicians justifiably relied on Defendant's  
7 representations and omissions.

8 135. Had appropriate warnings been provided, Plaintiff would not have  
9 consented to implantation.

10 136. As a direct and proximate result, Plaintiff suffered injuries including  
11 physical pain, emotional distress, surgical revision, and economic loss.

12 137. These claims are not preempted because they are based on parallel  
13 state-law duties that mirror federal requirements.

14 **WHEREFORE**, Plaintiff demands judgment against Defendant Boston  
15 Scientific Corporation for compensatory damages, attorneys' fees where permitted,  
16 costs of suit, interest, and all such further relief as the Court deems just and proper.

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18 **COUNT III – NEGLIGENCE PER SE AND BREACH OF FEDERAL**  
19 **REGULATORY DUTIES**

20 ***Against Defendant Boston Scientific***

21 (Florida Common Law; *Stengel v. Medtronic Inc.*, 704 F.3d 1224 (9th Cir. 2013);  
22 21 C.F.R. §§ 820.30(g), 820.75, 820.100, 820.198)

23 138. Plaintiff realleges and incorporates by reference all preceding  
24 paragraphs of this Complaint as though fully set forth herein.

25 139. At all relevant times, Defendant Boston Scientific Corporation owed  
26 duties to Plaintiff, her healthcare providers, and the public to comply with federal  
27 medical device regulations and to manufacture, monitor, and report on its spinal  
28

1 cord stimulator systems in accordance with Current Good Manufacturing  
2 Practices (“cGMPs”) set forth in 21 C.F.R. Part 820.

3 140. These cGMP requirements are not discretionary. They impose  
4 binding legal obligations to implement and maintain design controls, process  
5 validation, complaint handling systems, and corrective and preventive actions for  
6 Class III medical devices.

7 141. Defendant breached these duties by failing to:

- 8 • Implement effective design validation under 21 C.F.R. §  
9 820.30(g);
- 10 • Validate and monitor production processes under 21 C.F.R. §  
11 820.75;
- 12 • Respond adequately to known product failures through its  
13 Corrective and Preventive Action (CAPA) system in violation of 21  
14 C.F.R. § 820.100;
- 15 • Investigate and act upon post-market complaints in accordance  
16 with 21 C.F.R. § 820.198.

17 142. These regulatory breaches materially contributed to the manufacture,  
18 release, and continued distribution of unsafe, unvalidated, and defectively  
19 designed spinal cord stimulator systems that failed during normal use, including  
20 the Precision Montage MRI system implanted in Plaintiff.

21 143. Plaintiff and her physicians were members of the class of individuals  
22 the cGMP regulations were intended to protect – users of Class III implanted  
23 medical devices.

24 144. The harms suffered by Plaintiff, including physical pain, device  
25 failure, revision surgery, and ongoing neurological complications, are of the  
26 precise type that the cGMP regulations are intended to prevent.

27 145. Defendant’s violations of mandatory federal regulations constitute  
28 negligence per se under Florida law.

1 146. As a direct and proximate result of these regulatory violations,  
2 Plaintiff suffered physical injuries, including permanent injuries, emotional  
3 distress, medical costs, diminished enjoyment of life, additional surgical  
4 intervention, and other compensable harms.

5 147. These claims arise under state-law duties that genuinely parallel  
6 federal obligations imposed by the FDCA and are not expressly or impliedly  
7 preempted under *Riegel*, *Buckman*, or 21 U.S.C. § 360k(a).

8 **WHEREFORE**, Plaintiff demands judgment against Defendant Boston  
9 Scientific Corporation for compensatory damages, attorneys' fees where permitted,  
10 costs of suit, pre- and post-judgment interest, and such other and further relief as  
11 the Court deems just and proper.

12 **COUNT IV – BREACH OF EXPRESS WARRANTY**

13 ***Against Defendant Boston Scientific***

14 (Fla. Stat. § 672.314)

15 148. Plaintiff realleges and incorporates by reference all preceding  
16 paragraphs of this Complaint as though fully set forth herein.

17 149. At all relevant times, Defendant Boston Scientific Corporation  
18 expressly warranted, through its labeling, advertising, marketing materials,  
19 website representations, sales representatives, and field support personnel, that  
20 its spinal cord stimulator systems, including the Precision Montage MRI system,  
21 were safe, effective, durable, supported by clinical data, and fit for the treatment  
22 of chronic pain conditions.

23 150. Due to the extensive direct communications between Boston  
24 Scientific personnel and Plaintiff, privity of contract existed between these two  
25 parties.

26 151. These express warranties included, but were not limited to,  
27 assurances that:  
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- The safety and effectiveness of the Precision Montage MRI was validated by clinical data or clinical evidence;
- The permanent Precision Montage MRI system would provide greater or equivalent pain relief compared to the temporary trial SCS;
- The Precision Montage MRI system’s multiwaveform stimulation technology provided superior pain relief without significant risk of adverse side effects;
- The device was durable and appropriately designed for long-term implantation and therapeutic use without significant risks of lead migration, neurological injury, battery failure, rapidly declining efficacy, or stimulation-induced autonomic dysfunction;
- The system had been adequately tested for safety and effectiveness consistent with FDA requirements.

152. Such express warranties were made to Plaintiff’s healthcare providers and directly to Plaintiff through Boston Scientific’s promotional materials and field representatives, including Wayne (last name unknown).

153. Plaintiff and his healthcare providers reasonably relied on these express warranties in choosing to proceed with implantation of the Precision Montage MRI system and in making ongoing treatment decisions.

154. Contrary to these express warranties, the Precision Montage MRI system implanted in Plaintiff was defective, unreasonably dangerous, and incapable of delivering the promised therapeutic benefits. It was also not validated by clinical data, as represented to Plaintiff and her physicians. Instead, it caused Plaintiff to suffer injuries including persistent ineffective pain relief, nerve damage, device failure, stimulation-related complications, and ultimately, significant physical and emotional harm.

1 155. Defendant's conduct described herein breached its express  
2 warranties under Florida law.

3 156. Under Florida law, a manufacturer may be liable for breach of  
4 warranty if a product fails to perform as expressly warranted and causes harm.

5 157. Defendant's breaches of express warranty were a direct and  
6 proximate cause of Plaintiff's injuries, including physical pain, mental anguish,  
7 loss of enjoyment of life, additional medical expenses, and financial loss.

8 158. Plaintiff's claims for breach of express warranty are based on  
9 independent state-law duties and contractual obligations and are not expressly or  
10 impliedly preempted under federal law.

11 WHEREFORE, Plaintiff demands judgment against Defendant Boston  
12 Scientific Corporation for compensatory damages, consequential and incidental  
13 damages, punitive damages where permitted, attorneys' fees where allowed, costs  
14 of suit, pre- and post-judgment interest, and such other and further relief as the  
15 Court deems just and proper.

16 **COUNT V – BREACH OF IMPLIED WARRANTIES**

17 *Against Defendant Boston Scientific*

18 (Fla. Stat. §§ 672.313 and 672.314)

19 159. Plaintiff realleges and incorporates by reference all preceding  
20 paragraphs of this Complaint as though fully set forth herein.

21 160. Boston Scientific knew of the use that the Precision Montage MRI  
22 system was intended for and impliedly warranted to Plaintiff and his healthcare  
23 providers that the system was of merchantable quality and safe for the use for  
24 which it was intended, specifically the permanent implantation into human beings  
25 for the purpose of treating and relieving chronic pain.

26 161. Boston Scientific knew that, by holding the Precision Montage MRI  
27 system out as being approved by the FDA under the Premarket Approval process,  
28 physicians and patients would understand that the quality of the system was of a

1 degree that was supported by clinical data and evidence of safety and  
2 effectiveness. Thus, Boston Scientific impliedly warranted to Plaintiff and his  
3 physicians that the system was supported by such evidence.

4 162. Plaintiff and his healthcare providers reasonably relied on these  
5 implied warranties in choosing to proceed with implantation of the Precision  
6 Montage MRI system and in making ongoing treatment decisions.

7 163. Contrary to these implied warranties, the Precision Montage MRI  
8 system was not of merchantable quality, safe for its intended use, or supported by  
9 clinical data or evidence of safety and effectiveness.

10 164. Defendant's breaches of implied warranties were a direct and  
11 proximate cause of Plaintiff's injuries, including physical pain, mental anguish,  
12 loss of enjoyment of life, additional medical expenses, and financial loss.

13 165. Plaintiff's claims for breach of implied warranties are based on  
14 independent state-law duties and contractual obligations and are not expressly or  
15 impliedly preempted under federal law.

16 WHEREFORE, Plaintiff demands judgment against Defendant Boston  
17 Scientific Corporation for compensatory damages, consequential and incidental  
18 damages, punitive damages where permitted, attorneys' fees where allowed, costs  
19 of suit, pre- and post-judgment interest, and such other and further relief as the  
20 Court deems just and proper.

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22 **COUNT VI – FRAUDULENT MISREPRESENTATION**

23 ***Against Defendant Boston Scientific***

24 (Florida Commonlaw)

25 166. Plaintiff realleges and incorporates by reference all preceding  
26 paragraphs of this Complaint as though fully set forth herein.

27 167. At all relevant times, Defendant Boston Scientific Corporation, acting  
28 individually and through its agents, employees, and representatives, made

1 material misrepresentations and omissions of fact regarding the safety, efficacy,  
2 and regulatory compliance of its spinal cord stimulator systems, including the  
3 Precision Montage MRI system implanted in Plaintiff.

4 168. Defendant communicated directly with Plaintiff and her physicians  
5 through sales representatives, including Wayne and others.

6 169. Defendant represented to Plaintiff, her healthcare providers, and the  
7 general public, through marketing materials, sales presentations, Instructions for  
8 Use (IFU), direct to patient communications, and other communications, that:

- 9 • The Precision Montage MRI system was supported by clinical data  
10 and evidence of safety and effectiveness, as required for  
11 Premarket Approval, and was approved for sale based on such  
12 clinical data and evidence;
- 13 • The Precision Montage MRI system was safe and effective for the  
14 long-term management of chronic pain;
- 15 • The permanent Precision Montage MRI system would provide  
16 greater or equivalent pain relief compared to the temporary SCS  
17 trial;
- 18 • The device's multiwaveform stimulation capabilities reduced the  
19 risk of therapy failure and side effects compared to older systems;
- 20 • The device had been adequately validated and reviewed consistent  
21 with FDA requirements for significant design and functionality  
22 changes.

23 170. These material representations were false. Defendant knew or should  
24 have known, through reasonable investigation, post-market surveillance, and  
25 adverse event reporting, that:

- 26 • The Precision Montage MRI system was not supported by clinical  
27 data or evidence of safety and effectiveness, as required for  
28 Premarket Approval, nor was it approved for sale based on such  
clinical data or evidence;

- 1 • The Precision Montage MRI system was associated with a
- 2 materially increased risk of lead migration, therapy abandonment,
- 3 stimulation-induced autonomic dysfunction, device-related
- 4 complications, and rapid decrease in efficacy over time;
- 5 • The permanent device materially differed in performance and risk
- 6 profile from the external trial device;
- 7 • Boston Scientific had not submitted adequate clinical evidence to
- 8 substantiate the safety and effectiveness of its multiwaveform and
- 9 posture-adaptive programming technologies;
- 10 • Boston Scientific's devices were being improperly marketed under
- 11 serial PMA supplements rather than undergoing new PMA review
- 12 as required by law.

13 171. Defendant made these material misrepresentations and omissions

14 intentionally, willfully, recklessly, and with the intent to induce healthcare

15 providers to recommend, and patients to consent to, implantation of its devices.

16 172. Plaintiff's healthcare providers justifiably relied on Boston Scientific's

17 representations in recommending the Precision Montage MRI system, and

18 Plaintiff justifiably relied on the same representations in consenting to

19 implantation and subsequent medical decisions.

20 173. Boston Scientific's field representatives directly interacted with

21 Plaintiff and Plaintiff's care team and participated in programming the device.

22 Their conduct reinforced the impression that the system was safe and effective

23 and that post-implantation adjustments would resolve complications. These

24 communications omitted material facts and conveyed false assurances, including

25 before the implantation and for years after the implantation.

26 174. Plaintiff had no reasonable means of independently discovering the

27 falsity of Defendant's statements at the time of implantation, as Boston Scientific

28 had superior knowledge of the device's design changes, clinical performance, and

1 regulatory history, and actively concealed adverse information from physicians,  
2 patients, and regulators.

3 175. Plaintiff also had no reasonable means of independently discovering  
4 the falsity of Defendant's statements following implantation, as Boston Scientific  
5 had superior knowledge of the device's design changes, clinical performance, and  
6 regulatory history, and actively concealed adverse information from physicians,  
7 patients, and regulators.

8 176. Defendant's misrepresentations, including those made by sales  
9 representative Wayne, before the implantation procedure caused Plaintiff to  
10 consent to the implantation of the SCS system.

11 177. Defendant, through its representatives, also made misrepresentations  
12 to Plaintiff about complications and inadequate pain relief he experienced after  
13 the permanent implant, representing to her that multiple reprogrammings would  
14 alleviate her complications and increase her pain relief. These representations  
15 were made at the time of the reprogrammings.

16 178. Defendant knew these post-implant representations to be false.

17 179. Defendant's misrepresentations, including those made by Boston  
18 Scientific representatives in the months following the implantation procedure  
19 caused Plaintiff to keep the SCS system in his body long after it had any efficacy  
20 and after he began to experience complications, because Defendant's  
21 representatives assured him it would work with more adjustments and  
22 programming. This led to greater harm than he would have suffered if he had the  
23 SCS removed earlier.

24 180. Plaintiff discovered the probable causal relationship between his  
25 injuries and Defendant's conduct only after experiencing continued device-related  
26 complications and reviewing public disclosures, adverse event reports, and  
27 litigation materials that contradicted Defendant's original representations.  
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1 181. As a direct and proximate result of Defendant's fraudulent  
2 misrepresentations and omissions, Plaintiff suffered significant injuries, including  
3 physical pain, emotional distress, medical expenses, lost income, diminished  
4 quality of life, and other consequential damages.

5 182. Plaintiff's fraud claims arise under Florida common law, and are not  
6 based solely on FDCA violations, but rather Defendant's intentional  
7 misstatements and concealments made to induce reliance by Plaintiff and her  
8 physicians.

9 **WHEREFORE**, Plaintiff demands judgment against Defendant Boston  
10 Scientific Corporation for compensatory damages, punitive damages where  
11 permitted, attorneys' fees where allowed, costs of suit, pre- and post-judgment  
12 interest, and such other and further relief as the Court deems just and proper.

13 **COUNT VII – NEGLIGENT MISREPRESENTATION**

14 ***Against Boston Scientific***

15 (Florida Common Law)

16 183. Plaintiff realleges and incorporates by reference all preceding  
17 paragraphs of this Complaint as though fully set forth herein.

18 184. At all relevant times, Defendant Boston Scientific Corporation, acting  
19 individually and through its agents, employees, and representatives, owed a duty  
20 to exercise reasonable care in communicating truthful, accurate, and complete  
21 information about its spinal cord stimulator systems, including the Precision  
22 Montage MRI system.

23 185. Defendant supplied false information in its promotional materials,  
24 sales presentations, Instructions for Use (IFU), labeling, and direct  
25 communications to healthcare providers and patients regarding the safety,  
26 efficacy, design features, and regulatory status of the Precision Montage MRI  
27 system.

28 186. Specifically, Defendant negligently misrepresented that:

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- The safety and efficacy of the Precision Montage MRI system was validated by clinical data, as required for Premarket Approval, and was approved for sale based on such clinical data;
- The Precision Montage MRI system was safe and effective for the long-term treatment of chronic pain;
- The system’s multiwaveform stimulation would enhance therapeutic outcomes and reduce adverse effects compared to previous devices;
- The permanent Precision Montage MRI system would provide greater pain relief than the temporary SCS trial;
- The system had been appropriately validated through regulatory submissions consistent with FDA standards for safety and effectiveness.

187. Defendant made these representations without reasonable grounds for believing them to be true. A reasonable manufacturer exercising appropriate care would have known that:

- The Precision Montage MRI system was not supported by clinical data or evidence of safety and effectiveness, as required for Premarket Approval, nor was it approved for sale based on such clinical data or evidence;
- The Precision Montage MRI system was associated with a materially increased risk of lead migration, therapy abandonment, stimulation-induced autonomic dysfunction, and device-related complications;;
- The permanent implant differed materially in performance from the external trial device;
- The modified stimulation technologies and battery architecture introduced new and significant risks not present in predicate devices;

- 1 • Clinical data validating the long-term safety and efficacy of the
- 2 device modifications were lacking or insufficient;
- 3 • Adverse event trends, including stimulation-related autonomic
- 4 dysfunction, required disclosure to regulators and treating
- 5 physicians.

6 188. Plaintiff's healthcare providers reasonably relied on Defendant's  
7 representations in recommending implantation of the Precision Montage MRI  
8 system, and Plaintiff reasonably relied on the same representations in consenting  
9 to implantation.

10 189. Had Plaintiff and her healthcare providers been accurately and fully  
11 informed of the true risks and limitations associated with the device, they would  
12 not have elected to proceed with implantation or would have pursued alternative  
13 treatment options.

14 190. Defendant, through its representatives, also made misrepresentations  
15 to Plaintiff about complications and inadequate pain relief he experienced after  
16 the permanent implant, representing to her that multiple reprogrammings would  
17 alleviate her complications and increase her pain relief. These representations  
18 were made at the time of the reprogrammings.

19 191. Defendant knew or should have known that these post-implant  
20 representations were false.

21 192. Plaintiff relied on these representations in electing not to have the  
22 system removed earlier than he ultimately did, causing her additional and  
23 continued injuries.

24 193. As a direct and proximate result of Defendant's negligent  
25 misrepresentations, Plaintiff suffered physical injuries, emotional distress,  
26 additional medical expenses, financial losses, and diminished quality of life.

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1 194. Plaintiff’s negligent misrepresentation claims arise under state  
2 common law, including Florida Common Law. These claims are not premised  
3 solely on FDCA enforcement and are not preempted under *Buckman* or *Riegel*.

4 **WHEREFORE**, Plaintiff demands judgment against Defendant Boston  
5 Scientific Corporation for compensatory damages, attorneys’ fees where permitted,  
6 costs of suit, pre- and post-judgment interest, and such other and further relief as  
7 the Court deems just and proper.

8 **COUNT VIII – VIOLATIONS OF THE CALIFORNIA UNFAIR**  
9 **COMPETITION LAW**

10 ***Against Defendant Boston Scientific***

11 (Cal. Bus. & Prof. Code §§ 17200–17210; Cal. Civ. Code §§ 1750–1785)

12 195. Plaintiff realleges and incorporates by reference all preceding  
13 paragraphs of this Complaint as though fully set forth herein.

14 196. At all relevant times, Defendant Boston Scientific Corporation was  
15 engaged in trade and commerce as defined by the California Unfair Competition  
16 Law (“UCL”), including the nationwide marketing, sale, and distribution of spinal  
17 cord stimulator systems such as the Precision Montage MRI system.

18 197. The UCL prohibits unlawful, unfair, false, misleading, or deceptive  
19 acts and practices in the conduct of trade or commerce. Cal. Bus. & Prof. Code §  
20 17200.

21 198. Defendant Boston Scientific engaged in unlawful and deceptive  
22 conduct, including but not limited to:

- 23
- 24 • Misrepresenting the safety, reliability, and long-term performance  
25 of the Precision Montage MRI system;
  - 26 • Failing to disclose known risks of stimulation-induced autonomic  
27 dysfunction, device migration, and therapy failure;
  - 28 • Marketing the external trial device as predictive of permanent  
implant outcomes despite internal knowledge to the contrary;

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- Omitting material information regarding post-market failures, cGMP violations, and adverse events from its product labeling and promotional materials.

199. These acts and omissions were directed at Plaintiff, his healthcare providers, and the general public, and were likely to deceive reasonable consumers and physicians.

200. Boston Scientific’s conduct further violated the California Consumers Legal Remedies Act (“CLRA”), Cal. Civ. Code § 1770(a), including but not limited to:

- § 1770(a)(5): Representing that goods had characteristics or benefits which they did not have;
- § 1770(a)(7): Representing that goods were of a particular standard, quality, or grade when they were not;
- § 1770(a)(9): Advertising goods with the intent not to sell them as advertised;
- § 1770(a)(14): Representing that a transaction confers or involves rights or remedies it does not have.

201. Plaintiff and his healthcare providers reasonably relied on Boston Scientific’s misrepresentations and omissions in consenting to the device’s implantation and continuing therapy.

202. As a direct and proximate result, Plaintiff suffered injury in fact and economic damages, including unnecessary medical expenses, physical pain, emotional distress, and loss of quality of life.

203. Defendant’s violations of the UCL and CLRA were willful and knowing. Plaintiff seeks actual damages, restitution, injunctive relief, and statutory damages under Cal. Civ. Code § 1780, as well as civil penalties under Cal. Bus. & Prof. Code § 17206.



- 1 • Directly advised patients, including Plaintiff, about the safety and  
2 efficacy of the SCS system, benefits of the system, and impacts of  
3 reprogramming the SCS system;
- 4 • Directly advised patients, including Plaintiff, to keep the SCS  
5 system in their body despite complications and lack of efficacy of  
6 the system;
- 7 • Provided medical advice to patients, including Plaintiff, regarding  
8 the SCS system, symptoms now alleged to be caused by the SCS  
9 system, and decisions related to the SCS system.

10 207. These activities constitute the unlicensed practice of medicine in  
11 violation Florida law, which prohibit any person from diagnosing, treating, or  
12 recommending treatment for human ailments without a valid medical license.

13 208. Florida law permits statutes to serve as evidence of reasonableness,  
14 and therefore, violation of statute to serve as evidence of a breach of the duty to  
15 act reasonably toward others.

16 209. Plaintiff was a member of the class of individuals these statutes were  
17 intended to protect.

18 210. The harms suffered by Plaintiff, including physical pain, are the type  
19 that these statutes are intended to prevent.

20 211. As a direct and proximate result of Defendant's statutory violations  
21 and its failure to adequately train, supervise, or restrict the conduct of its  
22 representatives, Plaintiff suffered harm, including but not limited to: improper  
23 lead placement, ineffective stimulation, neurological injury, permanent nerve  
24 damage, device-related complications, additional surgeries, and prolonged pain  
25 and suffering.

26 212. Defendant's conduct constitutes negligence per se under Florida law  
27 and entitles Plaintiff to compensatory and punitive damages.  
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1 implanted in her body and undergo multiple reprogrammings, all while  
2 experiencing complications and poor pain relief.

3 219. The negligence of these representatives was willful and wanton,  
4 evidencing a conscious disregard for the rights of others and a desire to seek  
5 profits over patient safety. Therefore, punitive damages are appropriate.

6 220. Defendant Boston Scientific had a duty to reasonably and prudently  
7 hire, train, and supervise its employees, including the representatives that  
8 interacted with Plaintiff.

9 221. Defendant Boston Scientific breached its duty to reasonably and  
10 prudently hire, train, and supervise these employees.

11 222. If Defendant Boston Scientific had reasonably and prudently hired,  
12 trained, and supervised these employees, the representatives conduct described  
13 above would not have occurred and Plaintiff would not have been injured as a  
14 proximate result.

15 223. Therefore, Defendant Boston Scientific's failure to reasonably and  
16 prudently hire, train, and supervise these employees proximately caused  
17 Plaintiff's injuries.

18 **WHEREFORE**, Plaintiff respectfully requests that judgment be  
19 entered against Defendant Boston Scientific Corporation for compensatory and  
20 punitive damages, attorneys' fees where permitted, costs of suit, and all other  
21 relief the Court deems just and proper.

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1                    **COUNT XI – ADMINISTRATIVE PROCEDURE ACT (APA) –**  
2                    **DECLARATORY AND INJUNCTIVE RELIEF AGAINST THE FDA**

3                    ***Against Defendant U.S. Food and Drug Administration***

4                    (5 U.S.C. §§ 701–706; *Loper Bright Enterprises v. Raimondo*,  
5                    603 U.S. 369 (2024))

6                    224. Plaintiff realleges and incorporates by reference all preceding  
7 paragraphs of this Complaint as though fully set forth herein.

8                    225. This cause of action is brought against Defendant United States Food  
9 and Drug Administration solely in its official capacity under the Administrative  
10 Procedure Act, 5 U.S.C. §§ 701–706.

11                    226. The APA authorizes judicial review of final agency action, including  
12 agency actions that are arbitrary, capricious, an abuse of discretion, or otherwise  
13 not in accordance with law. See 5 U.S.C. §§ 706(1), 706(2)(A)–(D); *Loper Bright*  
14 *Enterprises v. Raimondo*, 603 U.S. 369 (2024).

15                    227. The FDA’s passive acceptance and approval of original PMA  
16 applications and subsequent supplements submitted by Defendant Boston  
17 Scientific Corporation for its Precision Montage MRI spinal cord stimulator  
18 system constituted final agency action within the meaning of the APA.

19                    228. The FDA acted arbitrarily and capriciously, and contrary to law, by:

- 20                    • Approving PMA P030017 despite Boston Scientific’s failure to supply  
21 data required by 21 CFR §814.20, and instead granting approval based  
22 on “available peer reviewed published literature for similar  
23 implantable spinal cord stimulation (SCS) systems.”
- 24                    • Approving substantial modifications to the original device –  
25 including the addition of multiwaveform stimulation, posture-  
26 adaptive programming, and battery redesign – without requiring  
27 new PMA applications or adequate independent clinical validation;  
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- 1 • Failing to adequately scrutinize Boston Scientific’s original PMA
- 2 Po30017 and subsequent PMA supplements that materially altered
- 3 the device’s design, safety profile, and intended use;
- 4 • Permitting Boston Scientific to evade full panel-track PMA review
- 5 through incremental supplement filings, despite knowing or having
- 6 reason to know that the cumulative changes were substantial;
- 7 • Failing to mandate labeling updates or field safety communications
- 8 in response to known adverse event trends related to stimulation-
- 9 induced autonomic dysfunction, device migration, and therapy
- 10 failure;
- 11 • Disregarding its statutory duty under 21 U.S.C. § 360e(d) to ensure
- 12 that significant changes to Class III devices receive the same rigorous
- 13 review as original PMA applications.

14 229. The FDA’s actions and omissions enabled Boston Scientific to market  
15 a materially altered, insufficiently validated, and defectively designed spinal cord  
16 stimulator system to Plaintiff and similarly situated patients without the  
17 protections mandated by Congress for high-risk medical device.

18 230. The FDA’s actions and omissions allowed Boston Scientific to market  
19 its SCS systems, including the Precision Montage MRI system, as Class III devices  
20 without subjecting these devices to the statutorily and regulatorily required  
21 review.

22 231. The FDA’s misconduct is further evidenced by its historical pattern of  
23 regulatory capture concerning spinal cord stimulators, including its decision to  
24 override an advisory panel recommendation in 2003 that implantable SCS devices  
25 be reclassified from Class III to Class II, without requiring manufacturers to  
26 complete PMA obligations thereafter. See FDA Docket No. 02P-0321.2

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27 <sup>2</sup> In 2001, Advanced Neuromodulation Systems (ANS) petitioned the FDA to reclassify  
28 implantable spinal cord stimulators from Class III to Class II. An FDA advisory panel  
recommended reclassification; however, the FDA headquarters overruled the panel and

1           232. Plaintiff suffered direct injury as a result of the FDA's arbitrary and  
2 unlawful agency actions. But for the FDA's approval of Boston Scientific's  
3 cumulative PMA supplement submissions without adequate scrutiny, Plaintiff  
4 would not have been implanted with the defective device that caused his injuries.

5           233. Plaintiff also suffered direct legal injury as a result of the FDA's  
6 arbitration and unlawful agency actions. But for the FDA's actions and omissions,  
7 Boston Scientific would not have received PMA approval for the SCS system  
8 implanted in Plaintiff, and therefore, would not be able to invoke federal  
9 preemption to shield itself from liability for the physical injuries that the SCS  
10 system caused Plaintiff.

11           234. The legal injury experienced by Plaintiff was a foreseeable result of  
12 the FDA's actions.

13           235. Plaintiff seeks declaratory relief declaring that the FDA's actions  
14 regarding the PMA supplements for Boston Scientific's spinal cord stimulator  
15 systems were arbitrary, capricious, an abuse of discretion, and contrary to law.

16           236. Plaintiff further seeks injunctive relief requiring the FDA to  
17 reconsider and, if necessary, rescind or suspend the PMA approvals granted for  
18 materially altered spinal cord stimulator systems that failed to undergo  
19 appropriate panel-track or original PMA review.

20           237. Plaintiff's claims under the APA are properly brought under 5 U.S.C.  
21 § 702 and 5 U.S.C. § 706, and are not precluded by any statutory or regulatory  
22 exemption from judicial review.

23           **WHEREFORE**, Plaintiff demands declaratory judgment against Defendant  
24 United States Food and Drug Administration, injunctive relief as permitted under  
25 the APA, costs of suit, reasonable attorneys' fees where permitted, and such other  
26 and further relief as the Court deems just and proper.

27 \_\_\_\_\_  
28 maintained Class III status without instituting corresponding PMA enforcement or  
strengthening requirements thereafter. *See* Docket No. 02P-0321 (FDA)

1 **VIII. PRAYER FOR RELIEF**

2 WHEREFORE, Plaintiff respectfully requests that this Court enter judgment  
3 in her favor and against Defendants Boston Scientific Corporation and the United  
4 States Food and Drug Administration, and award the following relief:

- 5 1. Compensatory damages for Plaintiff’s physical injuries, emotional  
6 distress, pain and suffering, loss of enjoyment of life, past and future  
7 medical expenses, and other economic and non-economic losses in an  
8 amount to be determined at trial;
- 9 2. Statutory damages and penalties as permitted under the California Unfair  
10 Competition Law and California Consumers Legal Remedies Act;
- 11 3. Consequential and incidental damages arising from Defendant’s breach of  
12 express and implied warranties under Florida law;
- 13 4. Punitive damages as permitted by Florida and California law for  
14 Defendant’s willful, fraudulent, and malicious conduct;
- 15 5. Declaratory relief pursuant to the Administrative Procedure Act declaring  
16 that the FDA’s actions in approving Boston Scientific’s PMA  
17 supplements without appropriate review were arbitrary, capricious, an  
18 abuse of discretion, and contrary to law;
- 19 6. Injunctive relief requiring the FDA to reconsider, rescind, or suspend  
20 PMA approvals for materially altered spinal cord stimulator devices that  
21 failed to undergo the statutorily required review processes;
- 22 7. Reasonable attorneys’ fees, costs of suit, and expenses incurred in this  
23 action as permitted by statute or common law;
- 24 8. Pre-judgment and post-judgment interest at the maximum rates permitted  
25 by law; and
- 26 9. Such other and further relief, whether at law or in equity, as this Court  
27 deems just and proper.  
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**JURY DEMAND**

Pursuant to Rule 38 of the Federal Rules of Civil Procedure, Plaintiff hereby demands a trial by jury on all issues so triable.

Dated: January 27, 2026

Respectfully submitted,

/s/ Trevor B. Rockstad  
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On Behalf of Brian Martini (Plaintiff)  
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