

**IN THE UNITED STATES DISTRICT COURT  
FOR THE SOUTHERN DISTRICT OF CALIFORNIA**

IN RE: ANGIODYNAMICS, INC.,  
AND NAVILYST MEDICAL, INC.,  
PORT CATHETER PRODUCTS  
LIABILITY LITIGATION

Case No.: 3:24-md-03125-JO-VET  
MDL No. 3125  
  
JUDGE JINSOOK OHTA

MALIA HOOKANO  
*Plaintiff,*  
  
vs.

COMPLAINT AND JURY DEMAND

ANGIODYNAMICS, INC., & NAVILYST  
MEDICAL, INC.,  
  
*Defendants.*

Civil Action No.:

**COMPLAINT**

Plaintiff files this Complaint pursuant to CMO No. 1, and is bound by the rights, protections, privileges, and obligations of that CMO. In accordance with CMO No. 1, Plaintiff hereby designates the United States District Court for the Central District of Nevada as Plaintiff’s venue as this case may have originally been filed there pursuant to 28 U.S.C. § 1391.

COMES NOW the Plaintiff, Malia Hookano, (who hereinafter shall be referred to as the “Plaintiff”), by and through her undersigned counsel, and brings this Complaint against AngioDynamics, Inc, and Navilyst Medical, Inc. (collectively, the “Defendants”), and alleges as follows:

1. This is an action for damages arising out of the failure relating to Defendants’ design, development, testing, assembling, manufacturing, packaging, promoting, marketing, distribution, supplying, and/or selling the defective

1 implantable vascular access device sold under the trade name of SmartPort  
2 (hereinafter “SmartPort”, or “Defective Device”).

3 **PARTIES**

4 2. Plaintiff, Malia Hookano, is an adult citizen of Henderson, Nevada.

5 3. Defendant AngioDynamics, Inc. (“AngioDynamics”) is a Delaware  
6 corporation with its principal place of business located in Latham, New York.  
7 AngioDynamics is engaged in the business of researching, developing, designing,  
8 licensing, manufacturing, distributing, supplying, selling, marketing, and  
9 introducing into interstate commerce, either directly or indirectly through third  
10 parties or related entities, its medical devices, including the SmartPort.

11 4. Defendant Navilyst Medical, Inc. (“Navilyst”) is a Delaware  
12 corporation with its principal place of business located in Marlborough,  
13 Massachusetts. Navilyst conducts business throughout the United States, including  
14 the State of Nevada, and is a wholly owned subsidiary of AngioDynamics. Navilyst  
15 is engaged in the business of researching, developing, designing, licensing,  
16 manufacturing, distributing, supplying, selling, marketing, and introducing into  
17 interstate commerce, either directly or indirectly through third parties or related  
18 entities, its medical devices, including the SmartPort.

19 **JURISDICTION AND VENUE**

20 5. This Court has subject matter jurisdiction over the parties pursuant to  
21 28 U.S.C. §1332(a) because the parties are citizens of different states and the  
22 amount in controversy exceeds \$75,000.00, exclusive of interest and cost.

23 6. Venue is proper in the Central District of Nevada pursuant to 28 U.S.C.  
24 §1391 by virtue of the facts that (a) a substantial part of the events or omissions  
25 giving rise to the claims occurred in that District and (b) Defendants’ products are  
26 produced, sold to and consumed by individuals in the State of Nevada, thereby  
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1 subjecting Defendants to personal jurisdiction in this action and making them all  
2 “residents” of that judicial District.

3 7. Defendants have and continue to conduct substantial business in the  
4 State of Nevada and in the Central District of Nevada, distribute vascular access  
5 products in that District, receive substantial compensation and profits from sales  
6 of vascular access products in that District, and made material omissions and  
7 misrepresentations and breaches of warranties in that District, so as to subject them  
8 to in personam jurisdiction in that District.

9 8. Consistent with the Due Process Clause of the Fifth and Fourteenth  
10 Amendments, both this Court and the Central District of Nevada have in personam  
11 jurisdiction over Defendants, because Defendants are present in the State of  
12 California and the State of Nevada, such that requiring an appearance does not  
13 offend traditional notions of fair and substantial justice.

14  
15 **PRODUCT BACKGROUND**

16 9. In or about 2007, a company called Rita Medical Systems, Inc.  
17 received clearance via the 510(k) Premarket Notification Program from the Food  
18 and Drug Administration (FDA) to market and sell a product called Vortex® CT  
19 Port Access System.

20 10. Around the same time Defendant AngioDynamics completed the  
21 acquisition of the assets and liabilities of Rita Medical Systems, Inc. and rebranded  
22 the subject product as SmartPort CT.

23 11. Defendants’ Vascular Access Devices were designed, patented,  
24 manufactured, labeled, marketed, sold, and distributed by the Defendants at all  
25 relevant times herein.

26 12. The SmartPort is one of several varieties of port/catheter systems that  
27 has been designed, manufactured, marketed, and sold by Defendants.

1 13. According to Defendants, the SmartPort is a totally implantable  
2 vascular access device designed to provide repeated access to the vascular system  
3 for the delivery of medication, intravenous fluids, parenteral nutrition solutions,  
4 and blood products.

5 14. The intended purpose of the SmartPort is to make it easier to deliver  
6 medications directly into the patient's bloodstream. The device is surgically placed  
7 completely under the skin and left implanted.

8 15. The SmartPort is a system consisting of two primary components: an  
9 injection port and a polyurethane catheter which includes additives intended to  
10 make it radiopaque.

11 16. The injection port has a raised center, or "septum," where the needle  
12 is inserted for delivery of the medication. The medication is carried from the port  
13 into the bloodstream through a small, flexible tube, called a catheter, that is inserted  
14 into a blood vessel.

15 17. The SmartPort is indicated for patient therapies requiring repeated  
16 access to the vascular system. The port system can be used for infusion of  
17 medications, I.V. fluids, parenteral nutrition solutions, blood products, and for the  
18 withdrawal of blood samples.

19 18. The product's catheter is marketed under the trade name Fluoromax  
20 and is comprised of a polymeric mixture of polyurethane, and a barium sulfate  
21 radiopacity agent. The Fluoromax catheter was first trademarked by Horizon  
22 Medical Products in 2005 and features a blue stripe which contains an even higher  
23 concentration of barium sulfate than the remainder of the lumen of the catheter.

24 19. Neither Horizon medical Products nor AngioDynamics received  
25 clearance from the FDA to market the Fluoromax catheter, making such device *per*  
26 *se* misbranded pursuant to the Food, Drug and Cosmetic Act.

1           20. Barium sulfate is known to contribute to reduction of the mechanical  
2 integrity of polyurethane *in vivo* as the particles of barium sulfate dissociate from  
3 the surface of the catheter over time, leaving microfractures and other alterations  
4 of the polymeric structure and degrading the mechanical properties of the  
5 polyurethane.

6           21. Researchers have shown that catheter surface degradation in products  
7 featuring a radiopaque barium sulfate stripe is concentrated at the locus of the  
8 stripe.<sup>1</sup>

9           22. The mechanical integrity of barium sulfate-impregnated polyurethane  
10 is affected by the concentration of barium sulfate as well as the heterogeneity of  
11 the modified polymer.

12           23. Upon information and belief, Defendants' manufacturing process in  
13 designing and constructing the catheter implanted in Plaintiff involved too high a  
14 concentration of barium sulfate particles for the polymer formulation, leading to  
15 improperly high viscosity of the admixed polyurethane before polymerization and  
16 causing improper mixing of barium sulfate particles within the polymer matrix.

17           24. This defect in the manufacturing process led to a heterogeneous  
18 modified polymer which led to an irregular catheter surface replete with fissure,  
19 pits and cracks.

20           25. Although the surface degradation and resultant mechanical failure can  
21 be reduced or avoided with design modifications (*e.g.*, using a higher grade  
22 radiopacity compound and/or encapsulating the admixed polymer within  
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27 <sup>1</sup> See Hecker JF, Scandrett LA. Roughness and thrombogenicity of the outer  
28 surfaces of intravascular catheters. *J Biomed Mater Res.* 1985;19(4):381-395.  
doi:10.1002/jbm.820190404

1 polyurethane), Defendants elected not to incorporate those design elements into  
2 the SmartPort.

3 26. At all times relevant, Defendants misrepresented the safety of the  
4 SmartPort system, and negligently designed, manufactured, prepared,  
5 compounded, assembled, processed, labeled, marketed, distributed, and sold the  
6 SmartPort system as safe and effective device to be surgically implanted to provide  
7 repeated access to the vascular system for the delivery of medications, intravenous  
8 fluids, parenteral nutrition solutions, and blood products.

9 27. At all times relevant to this action, Defendants knew and had reason  
10 to know, that the SmartPort was not safe for the patients for whom they were  
11 prescribed and implanted, because once implanted the device was prone to  
12 infection, thrombosis, fracturing, migrating, perforating internal vasculature and  
13 otherwise malfunctioning.

14 28. At all times relevant to this action, Defendants knew and had reason  
15 to know that patients implanted with a SmartPort port had an increased risk of  
16 suffering life threatening injuries, including but not limited to: death; hemorrhage;  
17 cardiac/pericardial tamponade (pressure caused by a collection of blood in the area  
18 around the heart); infection; cardiac arrhythmia and other symptoms similar to  
19 myocardial infarction; severe and persistent pain; and perforations of tissue,  
20 vessels and organs, or the need for additional surgeries to remove the defective  
21 device.

22 29. Soon after the SmartPort was introduced to market, which was years  
23 before Plaintiff was implanted with her device, Defendants began receiving large  
24 numbers of adverse event reports (“AERs”) from health care providers reporting  
25 that the SmartPort was fracturing post-implantation and that fractured pieces were  
26 migrating throughout the human body, including to the heart and lungs. Defendants  
27 also received large numbers of AERs reporting that the SmartPort was found to  
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1 have perforated internal vasculature. These failures were often associated with  
2 reports of severe patient injuries such as:

- 3 a. hemorrhage.
- 4 b. infection/ sepsis;
- 5 c. cardiac/pericardial tamponade;
- 6 d. cardiac arrhythmia and other symptoms similar to myocardial  
7 infarction;
- 8 e. severe and persistent pain;
- 9 f. perforations of tissue, vessels, and organs; and
- 10 g. upon information and belief, even death.

11 30. In addition to the large number of AERs which were known to  
12 Defendants and reflected in publicly accessible databases, there are many recorded  
13 device failures and/or injuries related to the Defendants' implantable port products  
14 which were concealed from medical professionals and patients through submission  
15 to the FDA's controversial Alternative Summary Reporting ("ASR") program.

16 31. The FDA halted the ASR program after its existence was exposed by  
17 a multi-part investigative piece, prompting a widespread outcry from medical  
18 professionals and patient advocacy groups.<sup>2</sup>

19 32. Prior to the discontinuation of the ASR program, Defendants reported  
20 numerous episodes of failures of their implanted port/catheter products – including  
21 numerous episodes of infection – under the ASR exemption, thereby concealing  
22 them from physicians and patients.

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27 <sup>2</sup> Christina Jewett, *Hidden Harm: Hidden FDA Reports Detail Harm Caused by*  
28 *Scores of Medical Devices*, Kaiser Health News (Mar. 2019).

1           33. Defendants were aware or should have been aware that the SmartPort  
2 had a substantially higher failure rate than other similar products on the market,  
3 yet Defendants failed to warn consumers of this fact.

4           34. Defendants also intentionally concealed the severity of complications  
5 caused by the SmartPort and the likelihood of these events occurring.

6           35. Rather than alter the design of the SmartPort to make it safer or  
7 adequately warn physicians of the dangers associated with the SmartPort,  
8 Defendants continued to actively and aggressively market the SmartPort as safe,  
9 despite their knowledge of numerous reports of infection, thrombosis, and  
10 associated injuries.

11           36. Moreover, Defendants concealed—and continue to conceal—their  
12 knowledge of the SmartPort’s dangerous propensity to precipitate infection.  
13 Defendants further concealed their knowledge that the catheter design caused these  
14 failures and that these failures cause serious injuries.

15           37. The conduct of Defendants, as alleged in this Complaint, constitutes  
16 willful, wanton, gross, and outrageous corporate conduct that demonstrates a  
17 conscious disregard for the safety of the Plaintiff. Defendants had actual  
18 knowledge of the dangers presented by the SmartPort, yet consciously failed to act  
19 reasonably to:

- 20           a. Adequately inform or warn Plaintiff, her prescribing physicians, or  
21           the public at large of these dangers;
- 22           b. Establish and maintain an adequate quality and post-market  
23           surveillance system; or
- 24           c. Recall the SmartPort System from the market.
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1                   **SPECIFIC FACTUAL ALLEGATIONS AS TO PLAINTIFF**

2           38. On or about February 4, 2019, Plaintiff underwent placement of an  
3   AngioDynamics SmartPort product at Mountainview Hospital, Nevada, by Muneer,  
4   Abdul MD. The device was implanted for the purpose of ongoing IV chemotherapy.

5           39. Defendants, directly or through their agents, apparent agents, servants,  
6   or employees designed, manufactured, marketed, advertised, distributed, and sold  
7   the SmartPort that was implanted in Plaintiff.

8           40. Defendants manufactured, sold, and/or distributed the SmartPort to  
9   Plaintiff, through her doctors, to be used for vein access.

10          41. On or about May 25, 2019, Plaintiff presented to N, Nevada, with  
11   complaints of fever, chills, and non-productive cough. Plaintiff was diagnosed with  
12   Sepsis and Pulmonary Embolism.

13          42. Defendants, directly or through their agents, apparent agents, servants,  
14   or employees designed, manufactured, marketed, advertised, distributed, and sold  
15   the SmartPort that was implanted in Plaintiff.

16          43. At all times, the SmartPort was utilized and implanted in a manner  
17   foreseeable to Defendants, as Defendants generated the instructions for use and  
18   created procedures for implanting the product.

19          44. The SmartPort implanted into Plaintiff was in the same or  
20   substantially similar condition as when it left the possession of Defendants, and in  
21   the condition directed by and expected by Defendants.

22          45. Plaintiff and her physicians foreseeably used and implanted the  
23   SmartPort, and did not misuse, or alter the SmartPort in an unforeseeable manner.

24          46. Defendants advertised, promoted, marketed, sold, and distributed the  
25   SmartPort as a safe medical device when Defendants knew or should have known  
26   the SmartPort was not safe for its intended purposes and that the product could  
27   cause serious medical problems.

1 47. Defendants had sole access to material facts concerning the defective  
2 nature of the products and their propensity to cause serious and dangerous side  
3 effects.

4 48. In reliance on Defendants' representations, Plaintiff's doctor was  
5 induced to, and did use the SmartPort.

6 49. As a result of having the SmartPort implanted, Plaintiff sustained  
7 significant mental and physical pain and suffering, suffered permanent injury,  
8 permanent and substantial physical deformity, underwent corrective surgery or  
9 surgeries, and suffered financial or economic loss, including, but not limited to,  
10 obligations for medical services and expenses.

11 50. Defendants' SmartPort was marketed to the medical community and  
12 to patients as safe, effective, reliable, medical devices; implanted by safe and  
13 effective, minimally invasive surgical techniques for the treatment of medical  
14 conditions, and as a safer and more effective as compared to the traditional  
15 products and procedures for treatment, and other competing Vascular Access  
16 Devices.

17 51. The Defendants have marketed and sold the SmartPort to the medical  
18 community at large and patients through carefully planned, multifaceted marketing  
19 campaigns and strategies. These campaigns and strategies include, but are not  
20 limited to, direct to consumer advertising, aggressive marketing to health care  
21 providers at medical conferences, hospitals, private offices, and/or group  
22 purchasing organizations, and include a provision of valuable consideration and  
23 benefits to the aforementioned.

24 52. The injuries, conditions, and complications suffered due to  
25 Defendants' SmartPort include but are not limited to hemorrhage;  
26 cardiac/pericardial tamponade; cardiac arrhythmia and other symptoms similar to  
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1 myocardial infarction; severe and persistent pain; perforations of tissue, vessels  
2 and organs; and even death.

3 53. Defendants were negligent toward Plaintiff in the following respects:

4 a. Defendant failed to design and establish a safe, effective procedure  
5 for removal of the SmartPort; therefore, in the event of a failure,  
6 injury, or complications, it is difficult to safely remove the SmartPort.

7 b. Defendants provided incomplete, insufficient, and misleading  
8 information to physicians in order to increase the number of  
9 physicians using the SmartPort for the purpose of increasing their  
10 sales. By so doing, Defendants caused the dissemination of  
11 inadequate and misleading information to patients, including the  
12 Plaintiff.

13 54. The SmartPort was utilized and implanted in a manner foreseeable to  
14 Defendants.

15 55. The SmartPort implanted into Plaintiff was in the same or  
16 substantially similar condition as when it left the possession of the Defendants, and  
17 in the condition directed by the Defendants.

18 56. At the time of her operation, Plaintiff was not informed of, and had  
19 no knowledge of the complaints, known complications, and risks associated with  
20 SmartPort.

21 57. Plaintiff was never informed by Defendants of the defective and  
22 dangerous nature of the SmartPort.

23 58. At the time of her implant, neither Plaintiff nor Plaintiff's physicians  
24 were aware of the defective and dangerous condition of the SmartPort.

25 59. At the time of the injuries referenced herein, Plaintiff did not know  
26 that the corrective surgery she underwent was due to a defect in the SmartPort.



1 67. Defendants' conduct, as described in this Complaint, also amounts to  
2 a continuing tort, and continues up through and including the date of the filing of  
3 Plaintiff's Complaint.

4 **FIRST CAUSE OF ACTION**

5 **NEGLIGENCE**

6 (Against Defendants AngioDynamics and Navilyst)

7 68. Plaintiff incorporates by reference the preceding paragraphs of this  
8 Complaint as if fully set forth herein.

9 69. The Defendants owed Plaintiff a duty to exercise reasonable care  
10 when designing, manufacturing, marketing, advertising, distributing, selling, and  
11 conducting post-market surveillance of the SmartPort.

12 70. The Defendants failed to exercise due care under the circumstances  
13 and therefore breached this duty by:

- 14 a. Failing to properly and thoroughly test the SmartPort before releasing  
15 the device to market, and/or failing to implement feasible safety  
16 improvements;
- 17 b. Failing to properly and thoroughly analyze the data resulting from any  
18 pre-market testing of the SmartPort;
- 19 c. Failing to conduct sufficient post-market testing and surveillance of  
20 the SmartPort;
- 21 d. Designing, manufacturing, marketing, advertising, distributing, and  
22 selling the SmartPort to consumers, including Plaintiff, without an  
23 adequate warning of the significant and dangerous risks of the  
24 SmartPort and without proper instructions to avoid the harm which  
25 could foreseeably occur as a result of using the SmartPort;
- 26 e. Failing to exercise due care when advertising and promoting the  
27 SmartPort; and  
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1 f. Negligently continuing to manufacture, market, advertise, and  
2 distribute the SmartPort after Defendants knew or should have known  
3 of its adverse effects.

4 71. As a direct and proximate result of the Defendants' actions, omissions  
5 and misrepresentations, Plaintiff was injured due to the use of the SmartPort, which  
6 caused Plaintiff various physical, mental, and emotional injuries and damages.

7 72. In performing the foregoing acts, omissions, and misrepresentations,  
8 Defendants acted grossly negligent, fraudulently, and with malice so as to justify  
9 an award of punitive and/or exemplary damages.

10 **SECOND CAUSE OF ACTION**

11 **STRICT PRODUCTS LIABILITY – FAILURE TO WARN**

12 (Against Defendants AngioDynamics and Navilyst)

13 73. Plaintiff incorporates by reference the preceding paragraphs of this  
14 Complaint as if fully set forth herein.

15 74. Defendants designed, set specifications, manufactured, prepared,  
16 compounded, assembled, processed, marketed, labeled, distributed, and sold the  
17 SmartPort, including the one implanted into Plaintiff, into the stream of commerce  
18 and in the course of same, directly advertised and marketed the device to  
19 consumers or persons responsible for consumers, and therefore had a duty to warn  
20 of the risk of harm associated with the use of the device and to provide adequate  
21 instructions on the safe and proper use of the device.

22 75. At the time Defendants designed, manufactured, prepared,  
23 compounded, assembled, processed, marketed, labeled, distributed, and sold the  
24 device into the stream of commerce, the device was defective and presented a  
25 substantial danger to users of the product when put to its intended and reasonably  
26 anticipated use, namely as an implanted port/catheter system to administer the  
27 medications. Defendants failed to adequately warn of the device's known or  
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1 reasonably scientifically knowable dangerous propensities, and further failed to  
2 adequately provide instructions on the safe and proper use of the device.

3 76. Defendants knew or should have known at the time they  
4 manufactured, labeled, distributed and sold the SmartPort that was implanted into  
5 Plaintiff that the SmartPort posed a significant and higher risk than other similar  
6 devices of device failure and resulting serious injuries.

7 77. Defendants failed to timely and reasonably warn of material facts  
8 regarding the safety and efficacy of the SmartPort; no reasonable health care  
9 provider, including Plaintiff's, or patient would have used the device in the manner  
10 directed, had those facts been made known to the prescribing healthcare providers  
11 or the consumers of the device.

12 78. The warnings, labels, and instructions provided by the Defendants at  
13 all times relevant to this action, are and were inaccurate, intentionally misleading,  
14 and misinformed and misrepresented the risks and benefits and lack of safety and  
15 efficacy associated with the device.

16 79. The health risks associated with the device as described herein are of  
17 such a nature that ordinary consumers would not have readily recognized the  
18 potential harm.

19 80. The device, which was designed, manufactured, prepared,  
20 compounded, assembled, processed, marketed, labeled, distributed, and sold into  
21 the stream of commerce by Defendants, was defective at the time of release into  
22 the stream of commerce due to inadequate warnings, labeling and/or instructions  
23 accompanying the product.

24 81. When Plaintiff was implanted with the device, Defendants failed to  
25 provide adequate warnings, instructions, or labels regarding the severity and extent  
26 of health risks posed by the device, as discussed herein.

1 82. Defendants intentionally underreported the number and nature of  
2 adverse events associated with dislodgement and migration of the devices to  
3 Plaintiff’s health care providers, as well as the FDA.

4 83. Neither Plaintiff nor her health care providers knew of the substantial  
5 danger associated with the intended and foreseeable use of the device as described  
6 herein.

7 84. Plaintiff and her health care providers used the SmartPort in a normal,  
8 customary, intended, and foreseeable manner, namely as a surgically placed device  
9 used to make it easier to deliver medications directly into the Plaintiff’s  
10 bloodstream. Moreover, Plaintiff’s health care providers did not place or maintain  
11 the device incorrectly such that it caused the device to “pinch off” or otherwise  
12 malfunction.

13 85. Upon information and belief, the defective and dangerous condition  
14 of the device, including the one implanted into Plaintiff, existed at the time they  
15 were manufactured, prepared, compounded, assembled, processed, marketed,  
16 labeled, distributed, and sold by Defendants to distributors and/or healthcare  
17 professionals or organizations. Upon information and belief, the device implanted  
18 in Plaintiff was in the same condition as when it was manufactured, inspected,  
19 marketed, labeled, promoted, distributed and sold by Defendants.

20 86. Defendants’ lack of sufficient warning and/or instructions was the  
21 direct and proximate cause of Plaintiff’s serious physical injuries, and economic  
22 damages in an amount to be determined at trial. In other words, had Defendants  
23 provided adequate warnings, Plaintiff and her physicians would not have used the  
24 device.

25 **THIRD CAUSE OF ACTION**

26 **STRICT PRODUCTS LIABILITY – DESIGN DEFECT**

27 (Against Defendants AngioDynamics and Navilyst)

1 87. Plaintiff incorporates by reference the preceding paragraphs of this  
2 Complaint as if fully set forth herein.

3 88. The SmartPort implanted in Plaintiff was not reasonably safe for its  
4 intended use and was defective with respect to its design.

5 89. The SmartPort was in a defective condition at the time that it left the  
6 possession or control of Defendants.

7 90. The SmartPort was unreasonably dangerous to the user or consumer.

8 91. The SmartPort was expected to and did reach the consumer without  
9 substantial change in its condition.

10 92. Defendants are strictly liable to the Plaintiff for designing,  
11 manufacturing, marketing, labeling, packaging and selling a defective product.

12 93. As a direct and proximate result of the SmartPort's aforementioned  
13 defects, Plaintiff was injured due to the use of the SmartPort, which caused Plaintiff  
14 various physical, mental, and emotional injuries and damages. Accordingly,  
15 Plaintiff seeks compensatory damages.

16 **FOURTH CAUSE OF ACTION**

17 **BREACH OF IMPLIED WARRANTY**

18 (Against Defendants AngioDynamics and Navilyst)

19 94. Plaintiff incorporates by reference the preceding paragraphs of this  
20 Complaint as if fully set forth herein.

21 95. Defendants impliedly warranted that the SmartPort was merchantable  
22 and fit for the ordinary purposes for which it was intended.

23 96. When the SmartPort was implanted in Plaintiff, it was being used for  
24 the ordinary purposes for which it was intended.

25 97. Plaintiff, individually and/or by and through her physician, relied  
26 upon Defendants' implied warranties of merchantability in consenting to have the  
27 SmartPort implanted in her.

1 98. Defendants breached these implied warranties of merchantability  
2 because the SmartPort implanted in Plaintiff was neither merchantable nor suited  
3 for its intended uses as warranted.

4 99. Defendants' breaches of their implied warranties resulted in the  
5 implantation of unreasonably dangerous and defective SmartPort in Plaintiff's  
6 body, placing said Plaintiff's health and safety in jeopardy.

7 100. The SmartPort was sold to Plaintiff's health care providers for  
8 implantation in patients, such as Plaintiff.

9 101. As a direct and proximate result of the SmartPort's aforementioned  
10 defects, Plaintiff was injured due to the use of the SmartPort, which caused Plaintiff  
11 various physical, mental, and emotional injuries and damages. Accordingly,  
12 Plaintiff seeks compensatory damages.

13 **FIFTH CAUSE OF ACTION**

14 **BREACH OF EXPRESS WARRANTY**

15 (Against Defendants AngioDynamics and Navilyst)

16 102. Plaintiff incorporates by reference the preceding paragraphs of this  
17 Complaint as if fully set forth herein.

18 103. Defendants through their officers, directors, agents, representatives,  
19 and written literature and packaging, and written and media advertisement,  
20 expressly warranted that the SmartPort was safe and fit for use by consumers, was  
21 of merchantable quality, did not produce dangerous side effects, and was  
22 adequately tested and fit for its intended use.

23 104. The SmartPort does not conform to the Defendants' express  
24 representations because it is not reasonably safe, has numerous serious side effects,  
25 and causes severe and permanent injury.

1 105. At all relevant times, the SmartPort did not perform as safely as an  
2 ordinary consumer would expect, when used as intended or in a reasonably  
3 foreseeable manner.

4 106. Plaintiff, her physicians, and the medical community reasonably  
5 relied upon the Defendants' express warranties for the SmartPort.

6 107. At all relevant times, the SmartPort was used on Plaintiff by Plaintiff's  
7 physicians for the purpose and in the manner intended by Defendants.

8 108. Plaintiff and Plaintiff's physicians, by the use of reasonable care,  
9 could not have discovered the breached warranty and realized its danger.

10 109. As a direct and proximate result of the SmartPort's aforementioned  
11 defects, Plaintiff was injured due to the use of the SmartPort, which caused Plaintiff  
12 various physical, mental, and emotional injuries and damages. Accordingly,  
13 Plaintiff seeks compensatory damages.

14 **SIXTH CAUSE OF ACTION**

15 **FRAUDULENT CONCEALMENT**

16 (Against Defendants AngioDynamics and Navilyst)

17 110. Plaintiff incorporates by reference the preceding paragraphs of this  
18 Complaint as if fully set forth herein.

19 111. Defendants fraudulently concealed information with respect to the  
20 SmartPort in the following particulars:

- 21 a. Defendants represented through the labeling, advertising, marketing  
22 materials, seminar presentations, publications, notice letters, and  
23 regulatory submissions that the SmartPort was safe and fraudulently  
24 withheld and concealed information about the substantial risks of  
25 using the SmartPort;
- 26 b. Defendants represented that the SmartPort was safer than other  
27 alternative systems and fraudulently concealed information which  
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1 demonstrated that the SmartPort was not safer than alternatives  
2 available on the market;

3 c. Defendants concealed that it knew these devices were fracturing and  
4 migrating from causes other than the manner in which the implanting  
5 physician implanted the device; and

6 d. That frequency of these failures and the severity of injuries were  
7 substantially worse than had been reported.

8 112. The Defendants had sole access to material facts concerning the  
9 dangers and unreasonable risks of the SmartPort.

10 113. The concealment of information by the Defendants about the risks of  
11 the SmartPort was intentional, and the representations made by Defendants were  
12 known by Defendants to be false.

13 114. The concealment of information and the misrepresentations about the  
14 SmartPort was made by the Defendants with the intent that Plaintiff's health care  
15 providers and Plaintiff rely upon them.

16 115. Plaintiff and her physicians relied upon the representations and were  
17 unaware of the substantial risks of the SmartPort which the Defendants concealed  
18 from the public, including Plaintiff and her physicians.

19 116. As a direct and proximate result of the SmartPort's aforementioned  
20 defects, Plaintiff was injured due to the use of the SmartPort, which caused Plaintiff  
21 various physical, mental, and emotional injuries and damages. Accordingly,  
22 Plaintiff seeks compensatory damages.

23 117. The Defendants acted with oppression, fraud, and malice towards  
24 Plaintiff, who accordingly requests that the trier of fact, in the exercise of its sound  
25 discretion, award additional damages for the sake of example and for the purpose  
26 of punishing Defendants for their conduct, in an amount sufficiently large to be an  
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1 example to others, and to deter this Defendants and others from engaging in similar  
2 conduct in the future.

3 118. Had Defendants not concealed this information, neither Plaintiff nor  
4 her health care providers would have consented to using the device in Plaintiff.

5 **SEVENTH CAUSE OF ACTION**

6 **NEVADA DECEPTIVE TRADE PRACTICES ACT**

7 (Against Defendants AngioDynamics and Navilyst)

8 119. Plaintiff incorporates by reference the preceding paragraphs of this  
9 Complaint as if fully set forth herein.

10 120. The acts and practices engaged in by Defendants constitute unlawful,  
11 unfair, deceptive, and/or fraudulent business or trade practices in violation of the  
12 Nevada Deceptive Trade Practices Act, (the “DTPA”), Deceptive Trade Practice  
13 Act NEV. REV. STAT. §§ 598.0915-598.0925.

14 121. Defendants engaged in unlawful practices, including unfair methods  
15 of competition, unconscionable acts or practices, and unfair or deceptive acts or  
16 practices in the conduct of trade or commerce in connection with the sale,  
17 distribution, and/or advertisement of the SmartPort in violation of the DTPA.

18 122. Defendants further engaged in unfair, unconscionable, deceptive,  
19 deliberately misleading, false, and/or deceptive acts and practices, all in violation  
20 of the DTPA, and as further described herein, including, but not limited to,  
21 misrepresenting that the SmartPort was reasonably safe for use and failing to  
22 adequately disclose the substantial risk of infection and harm the product entailed  
23 given the large number of adverse events Defendants knew or should have been  
24 aware of but did not adequately disclose to Plaintiff.

25 123. Defendants intended for Plaintiff, Plaintiff’s physicians, and other  
26 consumers to rely on their deceptive practices in order to continue selling and  
27 manufacturing the SmartPort.

1 124. Defendants' conduct was in or affecting commerce, namely,  
2 Defendants sold, and Plaintiff purchased the SmartPort, a product that Defendants  
3 falsely represented as having certain characteristics and benefits it did not have,  
4 inter alia, that it was reasonably safe for use, as further set forth above, in violation  
5 of the DTPA.

6 125. As a result of Defendants' conduct, Plaintiff suffered actual damages  
7 in that the product she purchased was misrepresented and worth far less than the  
8 product she thought she had purchased, had Defendants' representations been true.

9 **PUNITIVE DAMAGES**

10 126. Plaintiff is entitled to an award of punitive and exemplary damages  
11 based upon Defendants' intentional, willful, knowing, fraudulent, malicious acts,  
12 omissions, and conduct, and their complete and total reckless disregard for the  
13 public safety and welfare. Defendants intentionally and fraudulently  
14 misrepresented facts and information to both the healthcare community and the  
15 general public, including Plaintiff and her health care providers, by making  
16 intentionally false and fraudulent misrepresentations about the safety and efficacy  
17 of the SmartPort. Defendants intentionally concealed the true facts and information  
18 regarding the serious risks of harm associated with the implantation of said product,  
19 and intentionally downplayed the type, nature, and extent of the adverse side  
20 effects of being implanted with the device, despite Defendants' knowledge and  
21 awareness of the serious and permanent side effects and risks associated with use  
22 of same. Defendants further intentionally sought to mislead health care providers  
23 and patients, including Plaintiff and her health care providers, regarding the cause  
24 of infection and failures of the SmartPort.

25 127. Defendants had knowledge of and were in possession of evidence  
26 demonstrating that the SmartPort caused serious physical side effects. Defendants  
27 continued to market said product by providing false and misleading information  
28

1 with regard to the product’s safety and efficacy to the regulatory agencies, the  
2 medical community, and consumers of the SmartPort, notwithstanding Defendants’  
3 knowledge of the true serious side effects of the SmartPort, Defendants failed to  
4 provide accurate information and warnings to the healthcare community that would  
5 have dissuaded physicians from surgically implanting the SmartPort and  
6 consumers from agreeing to being implanted with the SmartPort, thus depriving  
7 physicians and consumers from weighing the true risks against the benefits of  
8 prescribing and implanting the SmartPort.

9 128. As a direct, proximate, and legal result of Defendants’ acts and  
10 omissions as described herein, and Plaintiff’s implantation with Defendants’  
11 defective product, Plaintiff suffered the injuries and damages described in this  
12 complaint.

13 **WHEREFORE**, Plaintiff demands judgment against Defendants for  
14 compensatory, special, and punitive damages, together with interest, costs of suit,  
15 attorneys’ fees, and all such other relief as the Court deems proper.

16 **PRAYER**

17 **WHEREFORE**, Plaintiff prays for judgment against each of the  
18 Defendants as follows:

- 19 a. Judgement be entered against all Defendants on all causes of action  
20 of this Complaint;
- 21 b. Plaintiff be awarded full, fair, and complete recovery for all claims  
22 and causes of action relevant to this action;
- 23 c. Plaintiff be awarded general damages according to proof at the time  
24 of trial;
- 25 d. Plaintiff be awarded damages, including past, present, and future,  
26 medical expenses according to proof at the time of trial;
- 27  
28

- e. Plaintiff be awarded punitive damages according to proof at the time of trial;
- f. Awarding pre-judgment and post-judgment interest to the Plaintiff;
- g. Awarding the costs and the expenses of this litigation to the Plaintiff; and
- h. For such other and further relief as the court may deem just and proper.

Respectfully submitted,

Dated: March 17, 2026

By: /s/ Gala Grand

Gala Grand, Esq.  
*Pro Hac Vice* Admission Pending  
**SEEGER WEISS LLP**  
 55 Challenger Road  
 Ridgefield Park, NJ 07660  
 Phone: (973) 639-5361  
 Fax: (973) 679-8656  
*Attorneys for Plaintiff*

**CERTIFICATE OF SERVICE**

I hereby certify that on March 17, 2026, a copy of the foregoing was served electronically and notice of the service of this document will be sent to all parties by operation of the Court’s electronic filing system to CM/ECF participants registered to receive service in this matter.

By: /s/ Gala Grand  
 Gala Grand, Esq.  
*Attorney for Plaintiff*

CIVIL COVER SHEET

The JS 44 civil cover sheet and the information contained herein neither replace nor supplement the filing and service of pleadings or other papers as required by law, except as provided by local rules of court. This form, approved by the Judicial Conference of the United States in September 1974, is required for the use of the Clerk of Court for the purpose of initiating the civil docket sheet. (SEE INSTRUCTIONS ON NEXT PAGE OF THIS FORM.)

I. (a) PLAINTIFFS
MALIA HOOKANO
(b) County of Residence of First Listed Plaintiff Clark County, NV
(c) Attorneys (Firm Name, Address, and Telephone Number) Gala Grand, Seeger Weiss, LLP, 55 Challenger Road, 6th Floor, Ridgefield Park, NJ 07660

DEFENDANTS
ANGIODYNAMICS, INC., ET AL.
County of Residence of First Listed Defendant Albany County, NY
NOTE: IN LAND CONDEMNATION CASES, USE THE LOCATION OF THE TRACT OF LAND INVOLVED.
Attorneys (If Known)
'26CV1682 JO VET

II. BASIS OF JURISDICTION (Place an "X" in One Box Only)
1 U.S. Government Plaintiff
2 U.S. Government Defendant
3 Federal Question (U.S. Government Not a Party)
4 Diversity (Indicate Citizenship of Parties in Item III)

III. CITIZENSHIP OF PRINCIPAL PARTIES (Place an "X" in One Box for Plaintiff and One Box for Defendant)
PTF DEF
Citizen of This State 1 1
Citizen of Another State 2 2
Citizen or Subject of a Foreign Country 3 3
Incorporated or Principal Place of Business In This State 4 4
Incorporated and Principal Place of Business In Another State 5 5
Foreign Nation 6 6

IV. NATURE OF SUIT (Place an "X" in One Box Only)
CONTRACT: 110 Insurance, 120 Marine, 130 Miller Act, 140 Negotiable Instrument, 150 Recovery of Overpayment & Enforcement of Judgment, 151 Medicare Act, 152 Recovery of Defaulted Student Loans (Excludes Veterans), 153 Recovery of Overpayment of Veteran's Benefits, 160 Stockholders' Suits, 190 Other Contract, 195 Contract Product Liability, 196 Franchise
REAL PROPERTY: 210 Land Condemnation, 220 Foreclosure, 230 Rent Lease & Ejectment, 240 Torts to Land, 245 Tort Product Liability, 290 All Other Real Property
TORTS: PERSONAL INJURY: 310 Airplane, 315 Airplane Product Liability, 320 Assault, Libel & Slander, 330 Federal Employers' Liability, 340 Marine, 345 Marine Product Liability, 350 Motor Vehicle, 355 Motor Vehicle Product Liability, 360 Other Personal Injury, 362 Personal Injury - Medical Malpractice
PERSONAL INJURY: 365 Personal Injury - Product Liability, 367 Health Care/Pharmaceutical Personal Injury Product Liability, 368 Asbestos Personal Injury Product Liability, 370 Other Fraud, 371 Truth in Lending, 380 Other Personal Property Damage, 385 Property Damage Product Liability
PRISONER PETITIONS: Habeas Corpus: 463 Alien Detainee, 510 Motions to Vacate Sentence, 530 General, 535 Death Penalty; Other: 540 Mandamus & Other, 550 Civil Rights, 555 Prison Condition, 560 Civil Detainee - Conditions of Confinement
FORFEITURE/PENALTY: 625 Drug Related Seizure of Property 21 USC 881, 690 Other
LABOR: 710 Fair Labor Standards Act, 720 Labor/Management Relations, 740 Railway Labor Act, 751 Family and Medical Leave Act, 790 Other Labor Litigation, 791 Employee Retirement Income Security Act
IMMIGRATION: 462 Naturalization Application, 465 Other Immigration Actions
BANKRUPTCY: 422 Appeal 28 USC 158, 423 Withdrawal 28 USC 157
INTELLECTUAL PROPERTY RIGHTS: 820 Copyrights, 830 Patent, 835 Patent - Abbreviated New Drug Application, 840 Trademark, 880 Defend Trade Secrets Act of 2016
SOCIAL SECURITY: 861 HIA (1395ff), 862 Black Lung (923), 863 DIWC/DIWW (405(g)), 864 SSID Title XVI, 865 RSI (405(g))
FEDERAL TAX SUITS: 870 Taxes (U.S. Plaintiff or Defendant), 871 IRS—Third Party 26 USC 7609
OTHER STATUTES: 375 False Claims Act, 376 Qui Tam (31 USC 3729(a)), 400 State Reapportionment, 410 Antitrust, 430 Banks and Banking, 450 Commerce, 460 Deportation, 470 Racketeer Influenced and Corrupt Organizations, 480 Consumer Credit (15 USC 1681 or 1692), 485 Telephone Consumer Protection Act, 490 Cable/Sat TV, 850 Securities/Commodities/Exchange, 890 Other Statutory Actions, 891 Agricultural Acts, 893 Environmental Matters, 895 Freedom of Information Act, 896 Arbitration, 899 Administrative Procedure Act/Review or Appeal of Agency Decision, 950 Constitutionality of State Statutes

V. ORIGIN (Place an "X" in One Box Only)
1 Original Proceeding, 2 Removed from State Court, 3 Remanded from Appellate Court, 4 Reinstated or Reopened, 5 Transferred from Another District (specify), 6 Multidistrict Litigation - Transfer, 8 Multidistrict Litigation - Direct File

VI. CAUSE OF ACTION
Cite the U.S. Civil Statute under which you are filing (Do not cite jurisdictional statutes unless diversity): 28 USC 1332 Diversity
Brief description of cause: Products Liability re: AngioDynamics Inc. and Navilyst Medical, Inc. Port Catheter Products

VII. REQUESTED IN COMPLAINT:
CHECK IF THIS IS A CLASS ACTION UNDER RULE 23, F.R.Cv.P. DEMAND \$ >\$75,000.00
CHECK YES only if demanded in complaint: JURY DEMAND: [X] Yes [ ] No

VIII. RELATED CASE(S) IF ANY (See instructions):
JUDGE Hon. Jinsook Ohta DOCKET NUMBER 3:24-md-03125-JO-VET

DATE 3/17/2026 SIGNATURE OF ATTORNEY OF RECORD /s/ Gala Grand

FOR OFFICE USE ONLY
RECEIPT # AMOUNT APPLYING IFP JUDGE MAG. JUDGE