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10		
11	ELIZABETH LANDERS, on behalf of her minor child, I.L.,	Case No.
12		
13	Plaintiff,	<u>COMPLAINT FOR</u>
14	V.	(1) Negligence
15	MERCK & CO., INC., and	 (2) Strict Liability (Failure to Warn) (2) Strict Liability (Manufacturing Defect)
16	MERCK SHARP & DOHME CORP,	(3) Strict Liability (Manufacturing Defect)(4) Breach of Warranty
17		(1) Dreach of Warranty (5) Common Law Fraud
18	Defendants.	(6) Violation of West Virginia's Consumer Credit and Protection Act
19		DEMAND FOR JURY TRIAL
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	COMPLAINT

1	COMES NOW Plaintiff, ELIZABETH LANDERS on behalf of her minor child I.L., who by
2	and through counsel, MORGAN & MORGAN, P.A., alleges against defendants MERCK & CO.,
3	INC., and MERCK, SHARP AND DOHME CORPORATION, and each of them, as follows:
4	INTRODUCTION
5	1. This common-law products liability, negligence, strict liability, breach of warranty and
6	fraud action arises out of serious and debilitating injuries, including but not limited to autonomic,
7	neurological and heterogenous autoimmune injuries and resulting sequalae that plaintiff, Elizabeth
8	Landers on behalf of her minor child I.L. ("Plaintiff"), sustained as a result of receiving injection of
9	the Gardasil vaccine, which was manufactured, labeled, and promoted by defendants Merck & Co.,
10	Inc., and Merck, Sharp and Dohme Corporation (collectively "Merck").
11	PARTIES AND VENUE
12	2. Plaintiff, I.L., represented by her mother Elizabeth Landers ("Plaintiff"), is a minor and
13	a resident and citizen of West Virginia.
14	3. Defendant Merck & Co., Inc., is a New Jersey corporation with its principal place of
15	business at One Merck Drive, Whitehouse Station, New Jersey.
16	4. Defendant Merck, Sharp and Dohme Corporation, is a New Jersey corporation with its
17	principal place of business at One Merck Drive, Whitehouse Station, New Jersey.
18	5. Defendants Merck & Co., Inc., and Merck, Sharp and Dohme Corporation shall
19	hereinafter collectively be referred to as "Merck."
20	6. At all times herein mentioned, each defendant was the agent, servant, partner, aider and
21	abettor, co-conspirator and/or joint venturer of the other defendants named herein and was at all times
22	operating and acting within the purpose and scope of said agency, service, employment, partnership,
23	conspiracy and/or joint venture and rendered substantial assistance and encouragement to the other
24	defendants, knowing that their collective conduct constituted a breach of duty owed to Plaintiff.
25	7. At all times herein mentioned, defendants were fully informed of the actions of their
26	agents and employees, and thereafter no officer, director or managing agent of defendants repudiated
27	those actions, which failure to repudiate constituted adoption and approval of said actions and all
28	defendants and each of them, thereby ratified those actions.

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8. There exists and, at all times herein mentioned there existed, a unity of interest in
 ownership between the named defendants, such that any individuality and separateness between the
 defendants has ceased and these defendants are the alter-ego of each other and exerted control over
 each other. Adherence to the fiction of the separate existence of these two named defendants as
 entities distinct from each other will permit an abuse of the corporate privilege and would sanction a
 fraud and/or would promote injustice.

9. At all times herein mentioned, the two Merck defendants were engaged in the business
of, or were successors in interest to, entities engaged in the business of researching, formulating,
compounding, testing, manufacturing, producing, processing, assembling, inspecting, distributing,
marketing, labeling, promoting, packaging, prescribing and/or advertising for sale, and selling
products for use by patients such as Plaintiff and her medical providers. As such, the two Merck
defendants are each individually, as well as jointly and severally, liable to Plaintiff for her damages.

10. The harm caused to Plaintiff resulted from the conduct of one or various combinations 13 of the two Merck defendants, and through no fault of Plaintiff. There may be uncertainty as to which 14 one or which combination of the two Merck defendants caused the harm. The two Merck defendants 15 have superior knowledge and information on the subject of which one or which combination of the 16 two defendants caused Plaintiff's injuries. Thus, the burden of proof should be upon each of the two 17 Merck defendants to prove that the defendant has not caused the harms Plaintiff has suffered. As 18 previously stated, the two named Merck defendants shall hereinafter and throughout this Complaint 19 be collectively referred to as "Merck." 20

11. Merck is the manufacturer, labeler and promoter of the Gardasil and Gardasil-9 21 vaccines, which are purported to be "cervical cancer vaccines" and "anal cancer vaccines" by 22 preventing a handful of the hundreds of strains of the Human Papillomavirus ("HPV"). Merck 23 regularly conducts and transacts business in West Virginia and has promoted Gardasil to consumers, 24 patients, hospitals, physicians, nurses and medical professionals, including but not limited to Plaintiff, 25 and the medical facility and medical professionals who prescribed and/or injected Plaintiff with 26 Gardasil. This Court has personal jurisdiction over Merck because defendants have sufficient 27 minimum contacts with West Virginia to render the exercise of jurisdiction by this Court proper. 28

1 12. This Court has subject matter jurisdiction over the parties pursuant to 28 U.S.C.
 2 §1332(a) because Plaintiff and the defendants are citizens of different states and the amount of
 3 controversy exceeds \$75,000.00, exclusive of interest and costs.

4 13. Venue is proper in this Court pursuant to 28 U.S.C. §1391 because a substantial portion
5 of the events and omissions giving rise to the claims asserted herein occurred in this District.

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I.

"History Doesn't Repeat Itself, But It Often Rhymes" – Mark Twain

GENERAL ALLEGATIONS

8 14. Merck traces its history back to 1668, when the original founder of the company,
9 Friedrich Jacob Merck, bought an apothecary in Darmstadt, Germany. The company operated as a
10 pharmacy for approximately the next 150+ years when, in 1827, Friedrich's descendant, Heinrich
11 Emmanuel Merck, converted the company into a drug manufacturing enterprise. Merck's first
12 products included morphine and cocaine.

13 15. Merck later manufactured a number of controversial products including Fosamax (a
14 purported bone density drug that caused bone fractures), Nuvaring (a birth control device associated
15 with life-threatening blood clots and death), and probably its most infamous drug, Vioxx (a pain
16 medication Merck was forced to pull from the market due to its cardiovascular risks), all of which
17 landed Merck in litigation hot water.

18 16. With regard to Vioxx, Merck was sued by tens of thousands of patients who alleged
19 they suffered heart attacks and other cardiovascular injuries as a result of ingesting the blockbuster
20 pain medication.

17. Documents unsealed during the Vioxx litigation in the early 2000s revealed a culture 21 wherein Merck knew early on that Vioxx was linked to fatal cardiovascular adverse events but 22 nonetheless intentionally chose to conceal these risks from the public and medical community and, 23 instead, orchestrated a scheme to downplay the severity of the risks. Merck misrepresented the results 24 of its clinical trials, failed to undertake the clinical trials that would reveal risks, and blacklisted 25 medical professionals who dared to publicly criticize the safety of Vioxx. See e.g., Eric J. Topol, 26 Failing the Public Health – Rofecoxib, Merck, and the FDA, 351 NEW ENGLAND JOURNAL OF 27 MEDICINE 1707 (2004); Gregory D. Curfman et al., Expression of Concern Reaffirmed, 354 NEW 28

ENGLAND JOURNAL OF MEDICINE 1193 (2006); Aaron S. Kesselheim et al., *Role of Litigation in Defining Drug Risks*, 17 JAMA 308 (2007); Harlan M. Krumholz et al., *What We Have Learnt From Vioxx*, 334 BRITISH MED. J. 120 (2007).

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18. The British Medical Journal reported that internal documents and communications 4 obtained from Merck during litigation revealed that Merck scientists internally acknowledged the 5 existence of Vioxx's risks very early on: "Since the early development of [Vioxx], some scientists at 6 Merck were concerned that the drug might adversely affect the cardiovascular system ... In internal 7 emails made public through litigation, Merck officials sought to soften the academic authors' 8 interpretation [of the data]. The academic authors changed the manuscript at Merck's request [to 9 make less of the apparent risk] ..." Harlan M. Krumholz et al., What We Have Learnt From Vioxx, 10 334 BRITISH MED. J. 120 (2007). And, despite Merck's knowledge of the risk, Merck never 11 conducted the necessary studies designed to evaluate cardiovascular risk. Id. 12

19. In an article published in the Journal of the American Medical Association, it was 13 reported that Merck worked to "diminish the impact of reported cardiovascular adverse effects by not 14 publishing adverse events and failing to include complete data on myocardial infarctions that occurred 15 during a key clinical trial. The information came to the public attention through a subpoena 5 years 16 after the article's publication, when [Vioxx] was already off the market." Aaron S. Kesselheim et al., 17 Role of Litigation in Defining Drug Risks, 17 JAMA 308 (2007). The article concludes: "These case 18 studies indicate that clinical trials and routine regulatory oversight as currently practiced often fail to 19 uncover important adverse effects for widely marketed products. In each instance, the litigation 20 process revealed new data on the incidence of adverse events, enabled reassessment of drug risks 21 through better evaluation of data, and influenced corporate and regulatory behavior." Id. 22

23 20. It was also revealed and reported that, in order to control the public narrative that Vioxx
24 was safe and risk free, "Merck issued a relentless series of publications...complemented by numerous
25 papers in peer-reviewed medical literature by Merck employees and their consultants. The company
26 sponsored countless continuing medical 'education' symposiums at national meetings in an effort to
27 debunk the concern about adverse cardiovascular effects." Eric J. Topol, *Failing the Public Health* –
28 *Rofecoxib, Merck, and the FDA*, 351 NEW ENGLAND JOURNAL OF MEDICINE 1707 (2004). In addition,

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Merck "selectively targeted doctors who raised questions about [Vioxx], going so far as pressuring
 some of them through department chairs." Harlan M. Krumholz et al., *What We Have Learnt From Vioxx*, 334 BRITISH MED. J. 120 (2007). Dr. Topol, Chairman of the Department of Cardiovascular
 Medicine at the Cleveland Clinic, commented: "Sadly, it is clear to me that Merck's commercial
 interest in [Vioxx] sales exceeded its concern about the drug's potential cardiovascular toxicity." Eric
 J. Topol, *Failing the Public Health – Rofecoxib, Merck, and the FDA*, 351 NEW ENGLAND JOURNAL
 OF MEDICINE 1707 (2004).

8 21. Once Merck's misdeeds vis-à-vis Vioxx were revealed in various jury trials, Merck paid
9 nearly \$5 billion to settle the tens of thousands of personal injury actions that had been brought
10 against it as a result of its concealment of Vioxx's cardiovascular risks. Merck paid an additional \$1
11 billion to settle a securities class action brought by investors who had lost money when Merck's stock
12 tanked following revelations of the drug's risks and subsequent lost sales. Merck was also forced to
13 pay \$950 million in civil and criminal fines to the Department of Justice and other governmental
14 entities as a result of various criminal activities Merck had engaged in with respect to Vioxx.

15 22. In 2005, Merck pulled Vioxx from the market and was desperate to find a replacement
16 for its previous multi-billion-dollar blockbuster.

17 23. Gardasil was viewed as the answer to the financial woes Merck had suffered from
18 Vioxx.

19 24. Indeed, some have euphemistically noted that HPV stood for "Help Pay for Vioxx."
20 25. In the aftermath of the Vioxx scandal, and seeking a replacement product, Merck's
21 senior director of clinical research, Eliav Barr, M.D., proclaimed of Gardasil: "This is it. *This is the*22 *Holy Grail!*"

II. In Bringing Its *Holy Grail*, Gardasil, to Market, Merck Engaged in the Same Fraudulent Research and Marketing It Had Engaged in Vis-à-vis Vioxx Resulting In Patients Being Exposed to a Vaccine That is Of Questionable Efficacy and Which Can Cause Serious and Debilitating Adverse Events

26 26. As outlined herein, in researching, developing, and marketing its new Holy Grail,
27 Gardasil, Merck engaged in the same unscrupulous tactics it had so infamously engaged in with
28 Vioxx.

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Certain Merck employees, scientists and executives involved in the Vioxx scandal were
 also involved with Gardasil, and it appears they employed the very same methods of manipulating
 science and obscuring risks as they did with Vioxx.

4 28. According to Merck's marketing claims, Gardasil (and, later, next-generation Gardasil
5 9) provided lifetime immunity to cervical, anal and other HPV-associated cancers.

6 29. As discussed more fully below, whether Gardasil prevents cancer (not to mention
7 lifetime immunity), is unproven. In fact, it may be more likely to cause cancer in those previously
8 exposed to HPV than to prevent it.

9 30. Moreover, Merck knows and actively conceals the fact that Gardasil can cause a
10 constellation of serious adverse reactions and gruesome diseases, including autoimmune diseases, and
11 death in some recipients.

31. As a result of Merck's fraud, Gardasil today is wreaking havoc on a substantial swath of
an entire generation of children and young adults on a worldwide scale.

14

A. Overview of the Human Papillomavirus

15 32. Human Papillomavirus ("HPV") is a viral infection that is passed between people
16 through skin-to-skin contact. There are more than 200 strains of HPV, and of those, more than 40
17 strains can be passed through sexual contact.

18 33. HPV is the most common sexually transmitted disease. It is so common that the
19 majority of sexually active people will get it at some point in their lives, even if they have few sexual
20 partners.

34. HPV, for the most part, is benign. More than 90 percent of HPV infections cause no
clinical symptoms, are self-limited, and are removed from the human body by its own immunological
mechanisms and disappear naturally from the body following an infection. *See, e.g.*, Antonio C. de
Freitas et al., *Susceptibility to cervical cancer: An Overview*, 126 GYNECOLOGIC ONCOLOGY 306
(August 2012).

35. Approximately 12 to 18 of the over 200 strains of HPV are believed to be associated
with cervical cancer, and approximately six of the strains are believed to be associated with anal
cancer.

1 36. Not every HPV infection puts one at risk for cervical cancer. Only persistent HPV 2 infections – not short-term or transient infections or sequential infections with different HPV types – in a limited number of cases with certain strains of the virus may cause the development of 3 precancerous lesions. With respect to cervical cancer, these precancerous lesions are typically 4 diagnosed through Pap smears and then removed through medical procedures. However, when 5 undiagnosed, they may in some cases progress to cervical cancer in some women. Other risk factors, 6 such as smoking, are also associated with cervical cancer. See Antonio C. de Freitas et al., 7 Susceptibility to cervical cancer: An Overview, 126 GYNECOLOGIC ONCOLOGY 305 (August 2012). 8 Infection with certain types of HPV are also associated with other diseases, such as genital warts. 9

37. Public health officials have long recommended the Pap test (also known as Pap Smear),
which detects abnormalities in cervical tissue, as the most effective frontline public health response to
the disease.

38. Since its introduction, cervical cancer screening through the Pap test has reduced the
rates of cervical cancer in developed countries by up to 80 percent. *Id.*

15 39. Incidences of cervical cancer have been declining dramatically worldwide as countries
16 have implemented Pap screening programs.

17 40. New cases of cervical cancer in the U.S. affect approximately 0.8 percent of women in
18 their lifetime. *See Cancer Stat Facts: Cervical Cancer*, NIH, at

https://seer.cancer.gov/statfacts/html/cervix.html. For those who are diagnosed, cervical cancer is
largely treatable, with a five-year survival rate of over 90 percent when the cancer is caught early. *See*Antonio C. de Freitas et al., *Susceptibility to cervical cancer: An Overview*, 126 GYNECOLOGIC
ONCOLOGY 305 (August 2012). Anal cancer is even more rare, and according to the current data,
approximately 0.2 percent of people will be diagnosed with anal cancer in their lifetime.

41. Although the incidence of cervical cancer was in rapid decline as a result of the
implementation of routine testing and screening, including the Pap test and various DNA testing
measures, Merck sought to fast-track a vaccine onto the market to prevent infection from four types of
HPV (only two of which are associated with cancer).

28

B. Overview of the Gardasil Vaccine and Its Fast-Tracked Approval

42. While there are over 200 types of the HPV virus, only 12 to 18 types currently are
 considered potentially associated with cervical or anal cancer. Merck's original Gardasil vaccine
 claimed to prevent infections from four strains (HPV Strain Types 6, 11, 16 and 18) and only two of
 those (Types 16 and 18) were associated with cervical and anal cancer.

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43. Under Food and Drug Administration ("FDA") requirements, to obtain approval for marketing a vaccine, the manufacturer must conduct studies to test the effectiveness and safety of the vaccine. Once FDA approval is obtained, the manufacturer has a duty to perform any further scientific and medical investigation as a reasonably prudent manufacturer would perform, and to engage in any necessary post-marketing pharmacovigilance related to the product.

44. The FDA approved Gardasil on June 8, 2006, after granting Merck fast-track status and
speeding the approval process to a six-month period, leaving unanswered material questions relating
to its effectiveness and safety as well as when and to whom the Gardasil vaccine ought to be
administered.

4 45. Merck failed, during the preapproval processing period and thereafter, to disclose (to
the FDA and/or the public), material facts and information relating to the effectiveness and safety of
Gardasil, as well as to whom the vaccine should or should not be administered.

Merck failed to perform in the preapproval processing period and thereafter, scientific
and medical investigations and studies relating to the safety, effectiveness and need for the Gardasil
vaccine as either required by and under FDA directives and regulations, and/or those which a prudent
manufacturer should have conducted unilaterally.

47. In June 2006, after the FDA's fast-tracked review, Gardasil was approved for use in
females ages nine through 26 for the purported prevention of cervical cancer and, almost immediately
thereafter, the Advisory Committee on Immunization Practices ("ACIP"), a committee within the
Centers for Disease Control ("CDC"), recommended Gardasil for routine vaccination of adolescent
girls ages eleven and twelve years old, but also allowed it to be administered to girls as young as nine
years old.

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48. On October 16, 2009, the FDA approved Gardasil for use in boys ages nine through 26

for the prevention of genital warts caused by HPV types 6 and 11, and in December 2010, it approved
 Gardasil for the purported prevention of anal cancer in males and females ages nine through 26.

49. Subsequently, Merck sought approval for Gardasil 9 (containing the same ingredients as
Gardasil, but in higher quantities), which purportedly guarded against five additional HPV strains
currently associated with cervical cancer and anal cancer (HPV Types 31, 33, 45, 52 and 58) than the
original Gardasil, for a total of nine strains.

50. The FDA approved Gardasil 9 in December 2014, for use in girls ages nine through 26
and boys ages nine through 15 for the purported prevention of cervical, vaginal, and anal cancers.
Presently, Gardasil 9 has been approved for and is being promoted by Merck to males and females
who are between nine and 45 years of age, with an emphasis by Merck on marketing to pre-teen
children and their parents. With little evidence of efficacy, the FDA also recently approved, on an
accelerated basis, Gardasil 9 for prevention of oropharyngeal and other head and neck cancers.

13 51. After the approval of the Gardasil 9 vaccine, the original Gardasil vaccine was phased
14 out of the U.S. Market; and the original Gardasil vaccine is no longer available for sale in the United
15 States.

16 52. According to data from the National Cancer Institute's ("NCI") Surveillance,
17 Epidemiology and End Results Program ("SEER"), the incidence of deaths from cervical cancer prior
18 to Gardasil's introduction in the United States had been steadily declining for years and, in 2006, was
19 2.4 per 100,000 women or approximately 1 in every 42,000 women. The currently available rate is
20 essentially unchanged, 2.2 per 100,000 women, based on data through 2017.

53. The median age of death from cervical cancer is 58, and death from anal cancer is 66,
and teenagers (who are the target population of Gardasil) essentially have zero risk of dying from
cervical or anal cancer.

54. Merck purchased fast-track review for Gardasil and Gardasil 9 under the Prescription
Drug User Fee Act ("PDUFA"). Fast-track is a process designed to facilitate the development of
drugs, and to expedite their review, in order to treat serious conditions and fill an unmet medical need.

27 55. Anxious to get Gardasil onto the market as soon as possible following the Vioxx
 28 debacle, Merck sought fast-track approval even though there already existed a highly effective and

side-effect free intervention, Pap smears, with no evidence that Gardasil was potentially superior to
 Pap smears in preventing cervical cancer.

56. In fact, the clinical trials Merck undertook did not even examine Gardasil's potential to 3 prevent cancer, rather, the trials only analyzed whether Gardasil could prevent potential precursor 4 conditions, i.e., HPV infections and cervical interepithelial neoplasia ("CIN") lesions graded from 5 CIN1 (least serious) to CIN3 (most serious), the vast majority of which resolve on their own without 6 intervention. CIN2 and CIN3 were the primary surrogate endpoints studied. Likewise, the clinical 7 trials from Gardasil did not examine Gardasil's potential to prevent anal cancer, rather, the trials 8 similarly only look at anal intraepithelial neoplasia ("AIN") lesions graded 1 through 3, and the 9 Gardasil 9 studies did not even include any studies concerning the efficacy of Gardasil in preventing 10 anal lesions. 11 57. According to the FDA, whether a condition is "serious" depends on such factors as 12 "survival, day-to-day functioning, or the likelihood that the condition, if left untreated, will progress 13 from a less severe condition to a more serious one." 14 As previously discussed, over 90 percent of HPV infections and the majority of cervical 58. 15 dysplasia, resolve without intervention. 16 59. However, Merck presented misleading data to the FDA suggesting that CIN2 and CIN3 17 inexorably result in cancer. 18 60. Federal law allows fast-track approval when there is no existing intervention to treat the 19 targeted disease or where the proposed treatment is potentially superior to an existing treatment. 20 61. Merck knows (and knew) that Gardasil and Gardasil 9 are far less effective than Pap 21 tests in preventing cervical cancer. 22 62. In order to obtain FDA approval, Merck designed and conducted a series of fraudulent 23 Gardasil studies and then influenced the votes of the FDA's Vaccines and Related Biological Products 24 Advisory Committee ("VRBPAC") and the CDC's Advisory Committee on Immunization Practices 25 ("ACIP") to win both an FDA license and a CDC/ACIP approval and recommendation that all 11 and 26 12 year old girls should be vaccinated with Gardasil. 27 63. That ACIP "recommendation" was, effectively, a mandate to doctors to sell Merck's 28

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very expensive vaccine, thereby compelling parents of American children as young as nine years old
 to buy this expensive product. With ACIP's recommendation, Merck was emboldened to build
 demand through direct-to-consumer advertising and door-to-door marketing to doctors, and, with the
 ACIP's blessing of the vaccine, circumvented the need to create a traditional market for the product.

64. Julie Gerberding, then the Director of CDC, obligingly ushered the Gardasil vaccine
through CDC's regulatory process manifestly ignoring clear evidence that Gardasil's efficacy was
unproven and that the vaccine was potentially dangerous.

8 65. Merck, shortly thereafter, rewarded Gerberding by naming her President of Merck
9 Vaccines in 2010.

66. In addition to the revolving regulatory/industry door, (wherein the Director of CDC 10 who approved the vaccine is subsequently employed by the manufacturer as a high-level executive to 11 oversee the commercial success of the vaccine she previously approved), it is also worth noting some 12 of the other conflicts of interest that exist within governmental agencies in relation to the facts 13 surrounding Gardasil. Scientists from the National Institute of Health ("NIH"), which is a division of 14 the United States Department of Health and Human Services ("HHS"), discovered a method of 15 producing "virus-like-particles" ("VLPs") that made creation of the Gardasil vaccine possible. The 16 NIH scientists' method of producing VLPs was patented by the Office of Technology Transfer 17 ("OTT"), which is part of the NIH, and the licensing rights were sold to Merck (for manufacture of 18 Gardasil). Not only does the NIH (and, in effect, the HHS) receive royalties from sales of Gardasil, 19 but the scientists whose names appear on the vaccine patents can receive up to \$150,000 per year (in 20 perpetuity). Accordingly, the Gardasil patents have earned HHS, NIH and the scientists who invented 21 the technology millions of dollars in revenue. 22

67. Moreover, members of ACIP have been allowed to vote on vaccine recommendations
even if they have financial ties to drug companies developing similar vaccines. According to a 2000
U.S. House of Representatives investigation report, the majority of the CDC's eight ACIP committee
members had conflicts of interest. The Chairman of ACIP served on Merck's Immunization Advisory
Board and a number of the other ACIP members had received grants, salaries, or other forms of
remuneration from Merck

1 2

C. Merck Engaged in Disease Mongering and False Advertising to Enhance **Gardasil Sales**

68. Both prior to and after the approval of Gardasil, Merck engaged in unscrupulous 3 marketing tactics designed to overemphasize both the risks associated with HPV and the purported 4 efficacy of Gardasil to scare the public into agreeing to mass vaccinations of the Gardasil vaccine.

5 69. Prior to Merck's aggressive marketing campaign, there was no HPV public health 6 emergency in high-resource countries, such as the United States.

7 70. Most women had never heard of HPV. The NCI's 2005 Health Information National 8 Trends Survey ("HINTS") found that, among U.S. women 18 to 75 years old, only 40 percent had 9 heard of HPV. Among those who had heard of HPV, less than half knew of an association between 10 HPV and cervical cancer. Furthermore, only four percent knew that the vast majority of HPV 11 infections resolve without treatment.

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71. The stage was set for Merck to "educate" the public about HPV, cervical cancer, and Gardasil, all to Merck's advantage.

14 72. Merck preceded its rollout of Gardasil with years of expensive disease awareness 15 marketing. Merck ran "Tell Someone" commercials, designed to strike fear in people about HPV and 16 cervical cancer – even ominously warning that you could have HPV and not know it. The 17 commercials could not mention Gardasil, which had not yet been approved by FDA, but did include 18 Merck's logo and name. Critics of Merck's pre-approval advertising and promotion called it 19 "deceptive and dishonest." While Merck claims the promotion was part of public health education, 20 critics complained that this "education" was designed to sell Gardasil and build the market for the 21 vaccine. See Angela Zimm and Justin Blum, Merck Promotes Cervical Cancer Shot by Publicizing 22 Viral Cause, BLOOMBERG NEWS, May 26, 2006.

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73. A year before obtaining licensing for its vaccine, Merck engaged in a major offensive in 'disease branding" to create a market for its vaccine out of thin air. See Beth Herskovits, Brand of the 24 25 Year, PHARMEXEC.COM, February 1, 2007. http://www.pharmexec.com/brand-year-0

26 74. Merck also engaged in a relentless propaganda campaign aimed at frightening and 27 guilting parents who failed to inoculate their children with Gardasil.

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75. In addition to paid advertising, Merck worked with third parties to "seed" an obliging

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1 media with terrifying stories about cervical cancer in preparation for Merck's Gardasil launch.

2 76. Prior to the FDA's 2006 approval of Gardasil, the mainstream media – under direction
3 of Merck and its agents – dutifully reported alarming cervical cancer stories, accompanied by the
4 promotion of an auspicious vaccine.

77. Merck intended its campaign to create fear and panic and a public consensus that "good
mothers vaccinate" their children with Gardasil. According to Merck propagandists, the only choice
was to "get the vaccine immediately" or "risk cervical or anal cancer."

8 78. Merck aggressively and fraudulently concealed the risks of the vaccine in broadcast
9 materials and in propaganda that it disseminated in the United States.

10 79. Merck sold and falsely promoted Gardasil knowing that, if consumers were fully
11 informed about Gardasil's risks and dubious benefits, almost no one would have chosen to vaccinate.

12 80. Merck negligently and fraudulently deprived parents and children of their right to13 informed consent.

81. One of Merck's television campaigns, conducted in 2016, shamelessly used child actors 14 and actresses, implicitly dying of cancer, looking straight into the camera and asking their parents 15 whether or not they knew that the HPV vaccine could have protected them against the HPV virus that 16 caused them to develop their cancers. Each actor asked the following question: "Did you know? 17 Mom? Dad?" See "Mom, Dad, did you know?" commercial: https://www.ispot.tv/ad/Ap1V/know-18 hpv-hpv-vaccination. Merck spent \$41 million over two months on the campaign. The ads said 19 nothing about potential side effects. Merck also distributed pamphlets via U.S. mail to doctors ahead 20 of the ad's release to encourage them to share it with their patients: 21

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82. Merck's fraudulent message was that cervical cancer and anal cancer were real-life
 killers of young men and women, notwithstanding the fact that the average age for development of
 cervical cancer is 50 years old, average age of development of anal cancer is 60 years old and that the
 cancer is virtually nonexistent in men and women under 20.

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83. Other television marketing campaigns Merck launched falsely proclaimed that Gardasil was a "cervical cancer vaccine" and that any young girl vaccinated with Gardasil would become "one less" woman with cervical cancer. The "One Less" marketing campaign portrayed Gardasil as if there were no question as to the vaccine's efficacy in preventing cervical cancer, and it disclosed none of Gardasil's side effects.

84. Merck marketed Gardasil with the most aggressive campaign ever mounted to promote
a vaccine, spending more on Gardasil advertising than any previous vaccine advertising campaign.

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D. Merck Used Scare Tactics and Provided Financial Incentives to Legislatures to Attempt to make the Gardasil Vaccine Mandatory for All School Children

14 85. An ACIP recommendation of a vaccine, adopted by individual states, opens the door to
 15 mandates affecting as many as four million children annually.

86. With Gardasil costing \$360 for the original three-dose series (exclusive of the necessary
doctor's visits) and Gardasil 9 now priced at \$450 for two doses (again, not including the cost of
doctor's visits), Merck stood to earn billions of dollars per year, in the US alone, with little marketing
costs.

20 87. Prior to Gardasil's approval in 2006, Merck was already targeting political figures to aid
 21 in the passage of mandatory vaccination laws.

88. As early as 2004, a group called Women in Government ("WIG") started receiving
funding from Merck and other drug manufacturers who had a financial interest in the vaccine.

89. With the help of WIG, Merck aggressively lobbied legislators to mandate Gardasil to all
sixth-grade girls. See Michelle Mello et al., Pharmaceutical Companies' Role in State Vaccination
Policymaking: The Case of Human Papillomavirus Vaccination, 102 AMERICAN J PUBLIC HEALTH
893 (May 2012).

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90. Dr. Diane Harper, a medical doctor and scientist who was hired as a principal
 investigator on clinical trials for Gardasil gave an interview for an article on the HPV vaccines and
 WIG in 2007. Harper, who had been a major presenter at a WIG meeting in 2005, stated that "the
 Merck representative to WIG was strongly supporting the concept of mandates later in the WIG
 meetings and providing verbiage on which the legislators could base their proposals."

6 91. WIG was one of dozens of "pay to play" lobby groups that Merck mobilized to push
7 HPV vaccine mandates.

8 92. Another group, the National Association of County and City Health Officials
9 (NACCHO), was also pushing HPV vaccine mandates in all 50 states.

10 93. To that end, Merck made large contributions to political campaigns and legislative
11 organizations. By February 2007, 24 states and the District of Columbia had introduced mandate
12 legislation.

13 94. Several states passed laws allowing preteen children as young as age 12 to "consent" to
14 vaccination with an HPV vaccine without parental consent or knowledge.

95. One New York state county offered children free headphones and speakers to encourage
them to consent to the Gardasil vaccine. *See* Mary Holland *et al.*, THE HPV VACCINE ON TRIAL:
SEEKING JUSTICE FOR A GENERATION BETRAYED 131 (2018).

96. Merck funneled almost \$92 million to Maryland's Department of Health between 2012
and 2018 to promote Gardasil in Maryland schools, in a fraudulent campaign that paid school officials
to deliberately deceive children and parents into believing Gardasil was mandatary for school
attendance. Josh Mazer, *Maryland should be upfront about HPV vaccinations for children*, CAPITAL
GAZETTE, August 14, 2018, at https://www.capitalgazette.com/opinion/columns/ac-ce-column-mazer-2018/0814-story.html.

24

E. Merck Pushed Gardasil Using Trusted Doctors and Third-Party Front Groups

97. In order to mobilize "third-party credibility" to push Gardasil, Merck gave massive
donations to dozens of nonprofit groups to "educate" the public via "education grants." For example,
a disclaimer on American College of Obstetricians and Gynecologists' Immunization for Women

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website stated that "[t]his website is supported by an independent educational grant from Merck and
 Sanofi Pasteur US."

3 98. Merck offered influential doctors (also known as "key opinion leaders") \$4,500 for
4 every Gardasil lecture they gave.

99. Among the allegedly independent organizations Merck recruited to push Gardasil were
the Immunization Coalition, the Allegheny County Board of Health, the Eye and Ear Foundation, the
Jewish Healthcare Foundation, the American Dental Association, the American College of
Obstetricians and Gynecologists, and the American Cancer Society.

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F. Merck Has Systematically Misrepresented the Efficacy of Gardasil By Advertising that Gardasil Prevents Cervical Cancer When There Are No Clinical Studies to Support This False Claim

11 100. Merck faced a daunting problem in convincing regulators, doctors, and the public to
12 accept the Gardasil vaccine.

13 101. Merck recommends the vaccine for children aged 11 to 12 years old, to provide
14 protection against a disease that, in the United States, is not generally diagnosed until a median age of
15 50. Moreover, in those rare instances of death, the median age is 58.

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102. There are no studies proving that Gardasil prevents cancer.

17 103. Because it can take decades for a persistent HPV infection to proceed to development of
18 cervical or anal cancer, and because cervical and anal cancers are so rare, a true efficacy study would
19 require decades and likely hundreds of thousand – if not millions – of trial participants to demonstrate
20 that eliminating certain HPV infections would actually prevent the development of cervical and anal
21 cancer.

104. Merck did not want to invest the time or money necessary to perform testing that would
 prove that its vaccine actually worked to prevent cervical and anal cancer.

105. Instead, Merck persuaded regulators to allow it to use "surrogate endpoints" to support
 its theory that the HPV vaccines would be effective in preventing cervical and anal cancer.

106. The clinical trials therefore did not test whether HPV vaccines prevent cervical, anal or
 other cancers. Instead, Merck tested the vaccines against development of certain cervical lesions,

 $\frac{1}{28}$ which some researchers suspect are precursors to cancer, although the majority of these lesions – even

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the most serious – regress on their own. See, e.g., Jin Yingji et al., Use of Autoantibodies Against
 Tumor-Associated Antigens as Serum Biomarkers for Primary Screening of Cervical Cancer, 8
 ONCOTARGET 105425 (Dec. 1, 2017); Philip Castle et al., Impact of Improved Classification on the
 Association of Human Papillomavirus With Cervical Precancer, 171 AMERICAN JOURNAL OF
 EPIDEMIOLOGY 161 (Dec. 10, 2009); Karoliina Tainio et al., Clinical Course of Untreated Cervical
 Intraepithelial Neoplasia Grade 2 Under Active Surveillance: Systematic Review and Meta-Analysis,
 360 BRIT. MED. J. k499 (Jan. 16, 2018).

8 107. The Department of Health and Human Services (HHS), which oversees the FDA, and
9 which also stood to make millions of dollars on the vaccine from patent royalties, allowed the use of
10 Merck's proposed surrogate endpoints.

11 108. The surrogate endpoints chosen by Merck to test the efficacy of its HPV vaccine were
12 cervical and anal intraepithelial neoplasia (CIN) grades 2 and 3 and adenocarcinoma in situ.

13 109. Merck used these surrogate endpoints even though it knew that these precursor lesions
14 are common in young women under 25 and rarely progress to cancer.

15 110. At the time FDA approved the vaccine, Merck's research showed only that Gardasil
16 prevented certain lesions (the vast majority of which would have resolved on their own without
17 intervention) and genital warts – not cancer itself, and only for a few years at that.

18 111. The use of these surrogate endpoints allowed Merck to shorten the clinical trials to a
19 few years and gain regulatory approvals of the vaccines without any evidence the vaccines would
20 prevent cancer in the long run.

112. Merck's advertisements assert that the HPV vaccine prevents cervical cancer. For
example, in a presentation to medical doctors, Merck proclaimed: "Every year that increases in
coverage [of the vaccine] are delayed, another 4,400 women will go on to develop cervical cancer."
The presentation goes on to tell doctors that women who do not get the vaccine will go on to develop
cancer.

26 113. Merck's foundational theory that HPV alone causes cervical and anal cancer, while
 27 dogmatically asserted, is not proven.

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1 114. Research indicates that cervical and anal cancer is a multi-factor disease with persistent
 2 HPV infections seeming to play a role, along with many other environmental and genetic factors,
 3 including smoking cigarettes or exposure to other toxic smoke sources, long-term use of oral
 4 contraceptives, nutritional deficiencies, multiple births (especially beginning at an early age), obesity,
 5 inflammation, and other factors. Not all cervical and anal cancer is associated with HPV types in the
 6 vaccines and not all cervical and anal cancer is associated with HPV at all.

7 115. Despite the lack of proof, Merck claimed that Gardasil could eliminate cervical and anal
8 cancer and other HPV-associated cancers.

9 116. However, *Merck knows* that the Gardasil vaccines cannot eliminate all cervical and anal
10 cancer or any other cancer that may be associated with HPV.

11 117. Even assuming the Gardasil vaccine is effective in preventing infection from the four to
12 nine vaccine-targeted HPV types, the results may be short term, not guaranteed, and ignore the 200 or
13 more other types of HPV not targeted by the vaccine, and some of which already have been associated
14 with cancer.

15 118. Even assuming these vaccine-targets are the types solely responsible for 100 percent of
16 cervical and anal cancer – which they are not – the vaccines have not been followed long enough to
17 prove that Gardasil protects girls and boys from cancer that would strike them 40 years later.

18 119. Under Merck's hypothetical theory, the reduction of pre-cancerous lesions should
19 translate to fewer cases of cervical and anal cancer in 30 to 40 years.

20 120. Cervical and anal cancer takes decades to develop and there are no studies that prove
21 the Gardasil vaccines prevent cancer.

121. In January 2020, a study from the UK raised doubts about the validity of the clinical
trials in determining the vaccine's potential to prevent cervical cancer. The analysis, carried out by
researchers at Newcastle University and Queen Mary University of London, revealed many
methodological problems in the design of the Phase 2 and 3 trials, leading to uncertainty regarding
understanding the effectiveness of HPV vaccination. *See* Claire Rees et al., *Will HPV Vaccine Prevent Cancer*? J. OF THE ROYAL SOC. OF MED. 1-15 (2020).

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1 122. As Dr. Tom Jefferson of the Centre for Evidence-Based Medicine pointed out: "The
 2 reason for choosing vaccination against HPV was to prevent cancer but there's no clinical evidence to
 3 prove it will do that."

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123. Gardasil has never been proven to prevent cervical or any other kind of cancer.

5 124. Yet Merck has marketed the Gardasil vaccines as if there is no question regarding their
6 efficacy at preventing cervical and anal cancer. In reality, they are at best protective against only four
7 to nine of the over 200 strains of the human papillomavirus.

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G. The Gardasil Vaccines Contain Numerous Hazardous Ingredients, Including At Least One Ingredient Merck Failed to Disclose to Regulators and the Public

Gardasil Contains A Toxic Aluminum Adjuvant

10 125. To stimulate an enhanced immune response that allegedly *might possibly* last for 50
 11 years, Merck added to the Gardasil vaccine a particularly toxic aluminum-containing adjuvant –
 12 Amorphous Aluminum Hydroxyphosphate Sulfate ("AAHS").

i.

126. Aluminum is a potent neurotoxin that can result in very serious harm.

14 127. The original Gardasil vaccine contains 225 micrograms of AAHS and Gardasil 9
15 contains 500 micrograms of AAHS.

16 128. Federal law requires that manufacturers cannot add adjuvants to vaccines that have not
17 been proven safe. 21 C.F.R. § 610.15(a).

18 129. AAHS has never been proven safe. AAHS is a recent proprietary blend of aluminum
and other unknown ingredients developed by Merck and used in Merck vaccines, including Gardasil.
20 Prior vaccines have used a different aluminum formulation.

130. Peer-reviewed studies show that aluminum binds to non-vaccine proteins, including the
host's own proteins, or to latent viruses, triggering autoimmune and other serious conditions. See
Darja Kanduc, *Peptide Cross-reactivity: The Original Sin of Vaccines*, 4 FRONTIERS IN BIOSCIENCE
1393 (June 2012).

131. Aluminum, including AAHS, has been linked to scores of systemic side effects
including, but not limited to: impairing cognitive and motor function; inducing autoimmune
interactions; increasing blood brain barrier permeability; inducing macrophagic myofascitis in muscle;
blocking neuronal signaling; interrupting cell-to-cell communications; corrupting neuronal-glial

interactions; interfering with synaptic transmissions; altering enzyme function; impairing protein 1 2 function; fostering development of abnormal tau proteins; and altering DNA.

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ii. Merck Lied About a Secret DNA Adjuvant Contained in The **Gardasil Vaccines**

Merck has repeatedly concealed or incorrectly identified Gardasil ingredients to the 132. FDA and the public.

6 133. Merck lied both to the FDA and the public about including a secret and potentially hazardous ingredient, HPV LI-DNA fragments, in Gardasil. These DNA fragments could act as a 8 Toll-Like Receptor 9 ("TLR9") agonist – further adjuvanting the vaccine and making it more potent. 9 Merck used this hidden adjuvant to prolong the immunological effects of the vaccine, but illegally 10 omitted it from its list of substances and ingredients in the vaccine.

134. Dr. Sin Hang Lee has opined that, without adding the TLR9 agonist, Gardasil would not be immunogenic. The DNA fragments bound to the AAHS nanoparticles act as the TLR9 agonist in both Gardasil and Gardasil 9 vaccines, creating the strongest immune-boosting adjuvant in use in any vaccine.

15 135. On multiple occasions, Merck falsely represented to the FDA and others, including 16 regulators in other countries, that the Gardasil vaccine did not contain viral DNA, ignoring the DNA 17 fragments.

18 136. This DNA adjuvant is not approved by the FDA and Merck does not list it among the ingredients as federal law requires. See 21 C.F.R. § 610.61(o) (requiring that adjuvants be listed on biologics' labeling). Even if not an adjuvant, the DNA fragments should have been listed because they represent a safety issue. 21 C.F.R. §610.61(n).

137. It is unlawful for vaccine manufacturers to use an experimental and undisclosed adjuvant.

24 When independent scientists found DNA fragments in every Gardasil vial tested, from 138. 25 all over the world, Merck at first denied, and then finally admitted, the vaccine does indeed include 26 HPV L1-DNA fragments.

27 28 1 139. Tellingly, Merck entered into a business arrangement with Idera Pharmaceuticals in
 2 2006 to explore DNA adjuvants to further develop and commercialize Idera's toll-like receptors in
 3 Merck's vaccine program.

4 140. To this day, the Gardasil package inserts do not disclose that DNA fragments remain in5 the vaccine.

6 141. Dr. Lee also found HPV DNA fragments from the Gardasil vaccine in post-mortem
7 spleen and blood samples taken from a young girl who died following administration of the vaccine.
8 See Sin Hang Lee, Detection of Human Papillomavirus L1 Gene DNA Fragments in Postmortem
9 Blood and Spleen After Gardasil Vaccination—A Case Report, 3 ADVANCES IN BIOSCIENCE AND
10 BIOTECHNOLOGY 1214 (December 2018).

11 142. Those fragments appear to have played a role in the teenager's death.
12 143. The scientific literature suggests there are grave and little-understood risks attendant to

13 injecting DNA into the human body.

14

iii. Gardasil Contains Borax

15 144. Gardasil contains sodium borate (borax). Borax is a toxic chemical and may have long16 term toxic effects.

17 145. Merck has performed no studies to determine the impact of injecting borax into millions18 of young children or adults.

19 146. Sodium borate is known to have adverse effects on male reproductive systems in rats,
 20 mice, and dogs. Furthermore, borax causes increased fetal deaths, decreased fetal weight, and
 21 increased fetal malformations in rats, mice, and rabbits.

147. The European Chemical Agency requires a "DANGER!" warning on borax and states
that borax "may damage fertility or the unborn child."

24 148. The Material Safety Data Sheet ("MSDS") for sodium borate states that sodium borate
25 "[m]ay cause adverse reproductive effects" in humans.

26 149. The FDA has banned borax as a food additive in the United States, and yet allows
27 Merck to use it in the Gardasil vaccine without any proof of safety.

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iv. Gardasil Contains Polysorbate 80

1 150. Gardasil contains Polysorbate 80. 2 151. Polysorbate 80 crosses the blood-brain barrier. Polysorbate 80 is used in drugs to open up the blood brain barrier in order to allow the 3 152. active ingredients in a drug to reach the brain and to elicit the intended response. It acts as an 4 emulsifier for molecules like AAHS and aluminum, enabling those molecules to pass through resistive 5 cell membranes. 6 153. Polysorbate 80 is associated with many health injuries, including, anaphylaxis, 7 infertility and cardiac arrest. 8 154. Polysorbate 80 was implicated as a cause, possibly with other components, of 9 anaphylaxis in Gardasil recipients in a study in Australia. See Julia Brotherton et al., Anaphylaxis 10 Following Quadrivalent Human Papillomavirus Vaccination, 179 CANADIAN MEDICAL ASSOC. J. 525 11 (September 9, 2008). Merck never tested Polysorbate 80 for safety in vaccines. 12 v. **Gardasil Contains Genetically Modified Yeast** 13 Gardasil contains genetically modified yeast. 155. 14 Studies have linked yeast with autoimmune conditions. See, e.g., Maurizo Rinaldi et 156. 15 al., Anti-Saccharomyces Cerevisiae Autoantibodies in Autoimmune Diseases: from Bread Baking to 16 Autoimmunity, 45 CLINICAL REVIEWS IN ALLERGY AND IMMUNOLOGY 152 (October 2013). 17 Study participants with yeast allergies were excluded from Gardasil clinical trials. 157. 18 158. Merck has performed no studies to determine the safety of injecting yeast into millions 19 of children and young adults. 20 // 21 H. As it Did in Vioxx, In Designing and Conducting Its Clinical Trials for 22 Gardasil, Merck Concealed Risks to Falsely Enhance the Safety Profile of Gardasil 23 24 159. Merck engaged in wholesale fraud during its safety and efficacy clinical studies. 25 160. In order to obtain its Gardasil license, Merck designed its studies purposefully to 26 conceal adverse events and exaggerate efficacy. 27 161. Merck sold Gardasil to the public falsely claiming that pre-licensing safety tests proved 28 it to be effective and safe.

1 162. In fact, Merck's own pre-licensing studies showed Gardasil to be of doubtful efficacy
 2 and dangerous.

3 163. The dishonesty in the clinical tests has led many physicians to recommend the
4 vaccination, under false assumptions.

5 164. The clinical trials clearly demonstrated that the risks of both Gardasil and Gardasil 9
6 vastly outweigh any proven or theoretical benefits.

7 165. Merck deliberately designed the Gardasil protocols to conceal evidence of chronic
8 conditions such as autoimmune diseases, menstrual cycle problems and death associated with the
9 vaccine during the clinical studies.

10 166. Merck employed deceptive means to cover up injuries that study group participants
11 suffered.

In early 2018, Lars Jørgensen, M.D., Ph.D. and Professor Peter Gøtzsche, M.D. (then 167. 12 with the Nordic Cochrane Centre), and Professor Tom Jefferson, M.D., of the Centre for Evidence-13 Based Medicine, published a study indexing all known industry and non-industry HPV vaccine 14 clinical trials and were disturbed to find that regulators such as the FDA and EMA (European 15 Medicines Agency) assessed as little as half of all available clinical trial results when approving the 16 HPV vaccines. Lars Jørgensen et al., Index of the Human Papillomavirus (HPV) Vaccine Industry 17 Clinical Study Programmers and Non-Industry Funded Studies: a Necessary Basis to Address 18 Reporting Bias in a Systematic Review, 7 SYSTEMATIC REVIEWS (January 18, 2018). 19

168. Per the indexing study discussed above, Merck appears to have kept a number of its
clinical trial results secret. Moreover, it appears that Merck reported only those findings that support
its own agenda.

23 169. Three separate reviews of the Gardasil vaccine by the Cochrane Collaboration found
24 that the trial data were "largely inadequate."

25 170. According to Dr. Tom Jefferson, "HPV [vaccine] harms have not been properly
 26 studied."

In 2019, numerous medical professionals published an article in the British Medical
 Journal outlining the flaws and incomplete nature of the publications discussing Merck's Gardasil

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clinical trials. The authors issued a "call to action" for independent researchers to reanalyze or 1 2 "restore the reporting of multiple trials in Merck's clinical development program for quadrivalent human papillomavirus (HPV) vaccine (Gardasil) vaccine." Peter Doshi et al., Call to Action: RIAT 3 Restoration of Previously Unpublished Methodology in Gardasil Vaccine Trials, 346 BRIT. MED. J. 4 2865 (2019). The authors explained that the highly influential publications of these studies, which 5 formed the basis of Gardasil's FDA approval, "incompletely reported important methodological 6 details and inaccurately describe the formulation that the control arm received, necessitating 7 correction of the record." Id. The authors explained that, while the publications claimed the clinical 8 trials of Gardasil were "placebo-controlled," "participants in the control arm of these trials did not 9 receive an inert substance, such as saline injection. Instead, they received an injection containing 10 [AAHS], a proprietary adjuvant system that is used in Gardasil to boost immune response." Id. 11

172. The researchers further opined that "the choice of AAHS-containing controls
complicates the interpretation of efficacy and safety results in trials … We consider the omission in
journal articles, of any rationale for the selection of AAHS-containing control, to be a form of
incomplete reporting (of important methodological details) and believe the rationale must be reported.
We also consider that use of the term 'placebo' to describe an active comparator like AAHS
inaccurately describes the formulation that the control arm received, and constitutes an important error
that requires correction." *Id.*

19 173. The authors pointed out that Merck's conduct "raises ethical questions about trial
20 conduct as well" and that they and other scientists would need to review the Gardasil clinical trial raw
21 data, in order to be able to analyze the safety and adverse event profile of Gardasil meaningfully and
22 independently. *Id.*

23

i. Small Clinical Trials

174. Although nine to 12-year-olds are the primary target population for HPV vaccines,
Merck used only a small percentage of this age group in the clinical trials. Protocol 018 was the only
protocol comparing children receiving a vaccine to those who did not. In that study, Merck looked at
results of fewer than 1,000 children 12 and younger for a vaccine targeting billions of boys and girls

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in that age group over time. In Protocol 018, 364 girls and 332 boys (696 children) were in the
 vaccine cohort, while 199 girls and 173 boys (372 children) received a non-aluminum control.

3 175. The small size of this trial means that it was incapable of ascertaining all injuries that
4 could occur as a result of the vaccine.

5

ii. Merck Used a Highly Toxic "Placebo" to Mask Gardasil Injuries

6 176. Instead of comparing health outcomes among volunteers in the Gardasil study group to
7 health outcomes among volunteers receiving an inert placebo, Merck purposefully used a highly toxic
8 placebo as a control in order to conceal Gardasil's risks in all trials using comparators with the
9 exception of Protocol 018, where only 372 children received a non-saline placebo containing
10 everything in the vaccine except the adjuvant and antigen.

11 177. Comparing a new product against an inactive placebo provides an accurate picture of
12 the product's effects, both good and bad. The World Health Organization ("WHO") recognizes that
13 using a toxic comparator as a control (as Merck did here) creates a "methodological disadvantage."
14 WHO states that "it may be difficult or impossible to assess the safety" of a vaccine when there is no
15 true placebo.

16 178. Merck deliberately used toxic "placebos" in the control group, in order to mask harms
17 caused by Gardasil to the study group.

18 179. Instead of testing Gardasil against a control with a true inert placebo, Merck tested its
19 vaccine in almost all clinical trials against its highly neurotoxic aluminum adjuvant, AAHS.

20 180. Merck gave neurotoxic aluminum injections to approximately 10,000 girls and young
21 women participating in Gardasil trials, to conceal the dangers of Gardasil vaccines.

181. Merck never safety tested AAHS before injecting it into thousands of girls and young
women in the control groups and the girls and young women were not told they could receive an
aluminum "placebo." Merck told the girls that they would receive either the vaccine or a safe inert
placebo.

182. Merck violated rules and procedures governing clinical trials when it lied to the clinical
study volunteers, telling them that the placebo was an inert saline solution – when in reality the
placebo contained the highly neurotoxic aluminum adjuvant AAHS.

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1	183. AAHS provoked terrible injuries and deaths in a number of the study participants when
2	Merck illegally dosed the control group volunteers with AAHS.
3	184. Since the injuries in the Gardasil group were replicated in the AAHS control group, this
4	scheme allowed Merck to falsely conclude that Gardasil's safety profile was comparable to the
5	"placebo."
6	185. The scheme worked and enabled Merck to secure FDA licensing.
7	186. Merck lied to the FDA when it told public health officials that it had used a saline
8	placebo in Protocol 018.
9	187. There was no legitimate public health rationale for Merck's failure to use a true saline
10	placebo control in the original Gardasil clinical trials. At that time, no other vaccine was yet licensed
11	for the four HPV strains Gardasil was intended to prevent.
12	188. A small handful of girls in a subsequent Gardasil 9 trial group, may have received the
13	saline placebo, but only after they had already received three doses of Gardasil for the Gardasil 9 trial.
14	iii. Merck Used Exclusionary Criteria to Further Conceal Gardasil Risks
14 15	
	Risks
15	Risks 189. Merck also manipulated the Gardasil studies by excluding nearly half of the original
15 16	Risks 189. Merck also manipulated the Gardasil studies by excluding nearly half of the original recruits to avoid revealing the effects of the vaccine on vulnerable populations.
15 16 17	Risks 189. Merck also manipulated the Gardasil studies by excluding nearly half of the original recruits to avoid revealing the effects of the vaccine on vulnerable populations. 190. After recruiting thousands of volunteers to its study, Merck excluded all women who
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 15 16 17 18 19 20 21 22 23 24 25 	Risks 189. Merck also manipulated the Gardasil studies by excluding nearly half of the original recruits to avoid revealing the effects of the vaccine on vulnerable populations. 190. After recruiting thousands of volunteers to its study, Merck excluded all women who had admitted to vulnerabilities that might be aggravated by the vaccine, such as abnormal Pap tests or a history of immunological or nervous system disorders. 191. Women could also be excluded for "[a]ny condition which in the opinion of the investigator might interfere with the evaluation of the study objectives." 192. Merck's protocol had exclusion criteria for subjects with allergies to vaccine ingredients including aluminum (AAHS), yeast, and the select enzymes. For most of these ingredients, there are limited resources for the public to test for such allergies in advance of being vaccinated. 193. Merck excluded anyone with serious medical conditions from the Gardasil clinical

1 194. Merck sought to exclude from the study all subjects who might be part of any subgroup
 2 that would suffer injuries or adverse reactions to any of Gardasil's ingredients.

195. The study exclusion criteria are not listed as warnings on the package inserts and the
package insert for Gardasil only mentions an allergy to yeast or to a previous dose of Gardasil as a
contraindication, rather than an allergy to any other component. Nonetheless, for most of the
ingredients, it is almost impossible to determine if such an allergy exists prior to being vaccinated and
Merck does not recommend allergy testing before administering the vaccine.

8 196. Instead of testing the vaccine on a population representative of the cross-section of
9 humans who would receive the approved vaccine, Merck selected robust, super-healthy trial
10 participants, who did not reflect the general population, in order to mask injurious effects on all the
11 vulnerable subgroups that now receive the vaccine. Therefore, the population tested in the clinical
12 trials was a much less vulnerable population than the population now receiving Gardasil.

13 14

iv. Merck Deceived Regulators and The Public by Classifying Many Serious Adverse Events, Which Afflicted Nearly Half of All Study Participants, As Coincidences

15 197. Because Merck did not use a true placebo, determining which injuries were attributable
16 to the vaccine and which were attributable to unfortunate coincidence was entirely within the
17 discretion of Merck's paid researchers.

18 198. In order to cover up and conceal injuries from its experimental vaccine, Merck, during
 19 the Gardasil trials, employed a metric, "new medical conditions," that allowed the company to dismiss
 20 and fraudulently conceal infections, reproductive disorders, neurological symptoms, and autoimmune
 21 conditions, which affected a troubling 50 percent of all clinical trial participants.

22 199. Merck's researchers systematically dismissed reports of serious adverse events from 49
23 percent of trial participants in order to mask the dangers of the vaccine.

24 200. Instead of reporting these injuries as "adverse events," Merck dismissed practically all
25 of these illnesses and injuries as unrelated to the vaccine by classifying them under its trashcan metric
26 "new medical conditions," a scheme Merck could get away with only because it used a "spiked"
27 (poisonous) placebo, that was yielding injuries at comparable rates.

28

201. Merck's use of a toxic placebo allowed the company to conceal from the public an
 epidemic of autoimmune diseases and other injuries and deaths associated with its multi-billion-dollar
 HPV vaccine.

202. Because Merck conducted its studies without a true placebo, Merck investigators had
wide discretion to decide what constituted an adverse event and used that power to dismiss a wave of
grave vaccine injuries, injuries that sickened half of the trial volunteers, as coincidental.

203. Almost half (49 percent) of all trial participants, regardless of whether they received the
vaccine or Merck's toxic placebo, reported adverse events, including serious illnesses such as blood,
lymphatic, cardiac, gastrointestinal, immune, musculoskeletal, reproductive, neurological and
psychological conditions, chronic illnesses such as thyroiditis, arthritis and multiple sclerosis, and
conditions requiring surgeries. *See, e.g.*, Nancy B. Miller, *Clinical Review of Biologics License Application for Human Papillomavirus 6, 11, 16, 18 L1 Virus Like Particle Vaccine (S. cerevisiae)*(*STN 125126 GARDASIL*), *manufactured by Merck, Inc.* at 393-94 (Table 302) (June 8, 2006).

14

15

v. Merck Manipulated the Study Protocols to Block Participants and Researchers from Reporting Injuries and Designed the Studies to Mask Any Long-Term Adverse Events

16 204. Merck adopted multiple strategies to discourage test subjects from reporting injuries.
17 205. Merck provided Vaccination Report Cards to a limited number of trial participants. For
18 example, in Protocol 015, only approximately 10 percent of participants – all in the United States,
19 despite trial sites worldwide – received Vaccination Report Cards to memorialize reactions in the first
20 few days following injections.

21 206. Furthermore, the report cards only included <u>categories</u> of "Approved Injuries" mainly
 22 jab site reactions (burning, itching, redness, bruising) leaving no room to report more serious
 23 unexplained injuries such as autoimmune diseases. In fact, they were designed for the purposes of
 24 reporting non-serious reactions only.

25 207. Furthermore, Merck instructed those participants to record information for only 14 days
26 following the injection.

27 208. In this way, Merck foreclosed reporting injuries with longer incubation periods or
28 delayed diagnostic horizons.

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209. Abbreviated reporting periods were part of Merck's deliberate scheme to conceal
 chronic conditions such as autoimmune or menstrual cycle problems, and premature ovarian failure,
 all of which have been widely associated with the vaccine, but would be unlikely to show up in the
 first 14 days following injection.

5 210. Merck researchers did not systematically collect adverse event data, from the trials,
6 which were spread out over hundreds of test sites all over the world.

7 211. To conceal the dangerous side effects of its vaccine, Merck purposely did not follow up
8 with girls who experienced serious adverse events during the Gardasil clinical trials.

9 212. Merck failed to provide the trial subjects a standardized questionnaire checklist of
10 symptoms, to document a comparison of pre- and post-inoculation symptoms.

11 213. To discourage its clinicians from reporting adverse events, Merck made the paperwork
 12 reporting requirements for supervising clinicians, onerous and time-consuming, and refused to pay
 13 investigators additional compensation for filling out the paperwork.

14 214. Thus, Merck disincentivized researchers from reviewing participants' medical records
15 even when the participant developed a "serious medical condition that meets the criteria for serious
16 adverse experiences" as described in the protocol.

17 215. Merck granted extraordinary discretion to its researchers to determine what constituted
18 a reportable adverse event, while incentivizing them to report nothing and to dismiss all injuries as
19 unrelated to the vaccine.

20 216. Merck used subpar, subjective data collection methods, relying on participants'
 21 recollections and the biased viewpoints of its trial investigators.

22 217. Merck downplayed the incidence of serious injuries and used statistical gimmickry to
 23 under-report entries.

24 218. During its Gardasil clinical trials, Merck failed to adequately capture and properly code
adverse events and symptoms, including but not limited to adverse events and symptoms that were
indicative of autoimmune or neurological injuries, including but not limited to POTS and CRPS, so as
to prevent the medical community, regulators and patients from learning about these adverse events

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and to avoid the responsibility of having to issue appropriate warnings concerning these adverse 1 2 events.

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Merck Deceived Regulators and the Public About Its Pivotal vi. Gardasil Clinical Trial (Protocol 018)

4 Merck tested Gardasil and Gardasil 9 in some 50 clinical trials, each one called a 219. "Protocol." However, results for many of these studies are not available to the public or even to the regulators licensing Gardasil. See Lars Jørgensen, et al., Index of the Human Papillomavirus (HPV) Vaccine Industry Clinical Study Programmers and Non-Industry Funded Studies: a Necessary Basis to Address Reporting Bias in a Systematic Review, 7 SYSTEMATIC REVIEWS 8 (January 18, 2018).

9 220. Gardasil's most important clinical trial was Protocol 018. The FDA considered 10 Protocol 018 the pivotal trial upon which Gardasil licensing approvals hinged, because FDA believed 11 1) it was the only trial where Merck used a "true saline placebo," and 2) it was the only trial with a 12 comparator group that included girls aged 11 to 12 – the target age for the Gardasil vaccine. See 13 Transcript of FDA Center For Biologics Evaluation And Research VRBPAC Meeting, May 18, 2006, 14 at 93 (Dr. Nancy Miller).

15 221. Merck lied to regulators, to the public and to subjects in its clinical trials by claiming 16 that the Protocol 018 "placebo" group received an actual saline or inert placebo.

17 222. When the FDA approved Gardasil, it described the Protocol 018 control as a "true 18 saline placebo."

19 The FDA declared that the Protocol 018 trial was "of particular interest" because Merck 223. 20 used a true saline placebo instead of the adjuvant as a control.

21 224. Merck told regulators that it gave a "saline placebo" to only one small group of 22 approximately 600 nine to 15-year-old children.

23 225. In fact, Merck did not give even this modest control group a true saline placebo, but 24 rather, the group members were given a shot containing "the carrier solution" – a witch's brew of 25 toxic substances including polysorbate 80, sodium borate (borax), genetically modified yeast, L-26 histidine, and possibly a fragmented DNA adjuvant.

27 The only components of Gardasil the control group did not receive were the HPV 226. 28 antigens and the aluminum adjuvant.

227. Despite the witches' brew of toxic chemicals in the carrier solution, those children fared
 much better than any other study or control group participants, all of whom received the AAHS
 aluminum adjuvant.

228. Only 29 percent of the vaccinated children and 31 percent of control recipients in
Protocol 018 reported new illnesses from Day 1 through Month 12, compared to an alarming 49.6
percent of those vaccinated and 49 percent of AAHS controls in the "pooled group" (composed of
some 10,000 young women and with the other participants combined) from Day 1 only through
Month 7 (not 12). Because the pooled group also included Protocol 018, even those numbers may not
be accurate with respect to those who received either a vaccine with a full dose of AAHS or those who
received an AAHS control.

11 229. Few of the participants in the Protocol 018 control group got systemic autoimmune
12 diseases, compared to 2.3 percent (1 in every 43) in the pooled group. In a follow-up clinical review
13 in 2008, the FDA identified three girls in the carrier-solution group with autoimmune disease. Based
14 on the number of girls in the placebo group as stated in the original 2006 clinical review, fewer than 1
15 percent of girls in the carrier solution group reported autoimmune disease.

16 230. In order to further deceive the public and regulators, upon information and belief,
17 Merck cut the dose of aluminum adjuvant in half when it administered the vaccine to the nine to
18 fifteen-year-old children in its Protocol 018 study group.

19 231. As a result, this group showed significantly lower "new medical conditions" compared
20 to other protocols.

21 232. Upon information and belief, Merck pretended that the vaccinated children in the
 22 Protocol 018 study group received the full dose adjuvant by obfuscating the change in formulation in
 23 the description.

24 233. Upon information and belief, Merck had cut the adjuvant in half, knowing that this
25 would artificially and fraudulently lower the number of adverse events and create the illusion that the
26 vaccine was safe.

234. Upon information and belief, Merck lied about this fact to the FDA.

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235. The data from that study therefore do not support the safety of the Gardasil formulation
 since Merck was not testing Gardasil but a far less toxic formulation.

3 236. Upon information and belief, Merck was testing a product with only half the dose of
4 Gardasil's most toxic component.

5 237. Upon information and belief, this is blatant scientific fraud, which continues to this day
6 because this is the study upon which current vaccine safety and long-term efficacy assurances are
7 based.

8 238. As set forth above, upon information and belief, Merck's deception served its purpose:
9 Only 29 percent of the vaccinated children in Protocol 018 reported new illness, compared to an
10 alarming 49.6 percent in the pooled group to receive the full dose adjuvant in the vaccine.

11

I. Contrary to Merck's Representations, Gardasil May Actually Cause and Increase the Risk of Cervical and Other Cancers

239. Gardasil's label states, "Gardasil has not been evaluated for potential to cause
carcinogenicity or genotoxicity." The Gardasil 9 label states: "GARDASIL9 has not been evaluated
for the potential to cause carcinogenicity, genotoxicity or impairment of male fertility.

15 240. Peer-reviewed studies, including CDC's own studies, have suggested that the
16 suppression of the HPV strains targeted by the Gardasil vaccine may actually open the ecological
17 niche for replacement by more virulent strains. *See* Fangjian Guo et al., *Comparison of HPV*

18 prevalence between HPV-vaccinated and non-vaccinated young adult women (20–26 years), 11

19 HUMAN VACCINES & IMMUNOTHERAPEUTICS 2337 (October 2015); Sonja Fischer et al., Shift in

20 prevalence of HPV types in cervical cytology specimens in the era of HPV vaccinations, 12

21 ONCOLOGY LETTERS 601 (2016); J. Lyons-Weiler, Biased Cochrane Report Ignores Flaws in HPV

22 Vaccine Studies, and Studies of HPV Type Replacement, (May 18, 2018). In other words, Gardasil

23 may increase the chances of getting cancer.

24 241. In short, the Gardasil vaccines, which Merck markets as anti-cancer products, may
25 themselves cause cancer or mutagenetic changes that can lead to cancer.

26 242. Merck concealed from the public data from its clinical trials indicating that the vaccines
27 enhance the risk of cervical cancers in many women.

28

243. Merck's study showed that women exposed to HPV before being vaccinated were 44.6
 percent more likely to develop cancerous lesions compared to unvaccinated women, even within a few
 years of receiving the vaccine.

4 244. In other words, Merck's studies suggest that its HPV vaccines may cause cancer in
5 women who have previously been exposed to HPV, particularly if they also have a current infection.

245. In some studies, more than 30 percent of girls show evidence of exposure to HPV
before age ten, from casual exposures, unwashed hands or in the birth canal. Flora Bacopoulou et al., *Genital HPV in Children and Adolescents: Does Sexual Activity Make a Difference?*, 29 JOURNAL OF
PEDIATRIC & ADOLESCENT GYNECOLOGY 228 (June 2016).

246. Even in light of the data demonstrating that Gardasil can increase the risk of cancer in
girls who previously have been exposed to HPV, in order to increase profits, Merck's Gardasil labels
and promotional material do not inform patients and medical doctors of this important risk factor.

247. Some clinical trial participants have developed cancer, including cervical cancer.

14 248. Numerous women have reported a sudden appearance of exceptionally aggressive15 cervical cancers following vaccination.

13

16 249. Cervical cancer rates are climbing rapidly in all the countries where Gardasil has a high
17 uptake.

18 250. An Alabama study shows that the counties with the highest Gardasil uptakes also had19 the highest cervical cancer rates.

20 251. After the introduction of HPV Vaccine in Britain, cervical cancer rates among young
21 women aged 25 to 29 has risen 54 percent.

22 252. In Australia, government data reveals there has been a sharp increase in cervical cancer
23 rates in young women following the implementation of the Gardasil vaccine. The most recent data
24 reveal that, 13 years after Gardasil was released and pushed upon teenagers and young adults, there
25 has been a 16 percent increase in 25 to 29 year-olds and a 30 percent increase in 30 to 34 year-old
26 girls contracting cervical cancer, corroborating the clinical trial data that Gardasil may *increase* the
27 risk of cervical cancer, particularly in patients who had previous HPV infections. Meanwhile, rates
28 are decreasing for older women (who have not been vaccinated).

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1	253. In addition to the belief that Gardasil may create and open an ecological niche for						
2	replacement by more virulent strains of HPV, resulting in the increase of cervical cancers as outlined						
3	above, in light of Merck's false advertising that Gardasil prevents cervical cancer, young women who						
4	have received Gardasil are foregoing regular screening and Pap tests in the mistaken belief that HPV						
5	vaccines have eliminated all their risks.						
6	254. Cervical screening is proven to reduce the cases of cervical cancer, and girls who have						
7	taken the vaccine are less likely to undergo cervical screenings.						
8	255. Data show that girls who received HPV vaccines before turning 21 are far less likely to						
9	get cervical cancer screening than those who receive the vaccines after turning 21.						
10	256. The cervical screening is more cost effective than vaccination alone or vaccination with						
11	screening.						
12	257. Therefore, Pap tests, which detect cervical tissue abnormalities, and HPV DNA testing						
13	are the most effective frontline public health response to cervical health.						
14 15	J. Merck has Concealed the Fact that Gardasil Induces and Increases the Risk of Autoimmune Diseases, and Other Injuries, Including But Not Limited to, Postural Orthostatic Tachycardia Syndrome, Chronic Fatigue Syndrome, Neuropathy, Fibromyalgia and Dysautonomia						
16	258. Gardasil induces and increases the risk of autoimmune disease.						
17	259. Gardasil has been linked to a myriad of autoimmune disorders, including but not						
18	limited, to: Guillain-Barré syndrome ("GBS"), postural orthostatic tachycardia syndrome ("POTS"),						
19	Orthostatic Intolerance ("OI"), chronic inflammatory demyelinating polyneuropathy ("CDIP"), small						
20	fiber neuropathy ("SNF"), systemic lupus erythematosus ("SLE"), immune thrombocytopenic						
21	purpura ("ITP"), multiple sclerosis ("MS"), acute disseminated encephalomyelitis ("ADEM"),						
22	antiphospholipid syndrome ("APS"), transverse myelitis, rheumatoid arthritis, interconnective tissue						
23	disorder, autoimmune pancreatitis ("AIP") and autoimmune hepatitis.						
24	260. Gardasil has also been linked to a myriad of diseases and symptoms that are associated						
25	with induced-autoimmune disease, including for example, fibromyalgia, dysautonomia, premature						
26	ovarian failure, chronic fatigue syndrome ("CFS"), chronic regional pain syndrome ("CRPS"),						
27	cognitive dysfunction, migraines, severe headaches, persistent gastrointestinal discomfort, widespread						
28							
	29						

pain of a neuropathic character, encephalitis syndrome, autonomic dysfunction, joint pain, and brain
 fog.

261. In a 2015 textbook, VACCINES AND AUTOIMMUNITY, edited by Dr. Yehuda Shoenfeld,
the father of autoimmunology research, and many of the world's leading autoimmunity experts, the
scientists concluded that Gardasil can cause autoimmune disorders because of the vaccine's strong
immune stimulating ingredients. *See* Lucija Tomljenovic & Christopher A. Shaw, *Adverse Reactions to Human Papillomavirus Vaccines*, VACCINES & AUTOIMMUNITY 163 (Yehuda Shoenfeld et al. eds.,
2015).

9 262. Medical experts have opined that the mixture of adjuvants contained in vaccines, in
10 particular in the Gardasil vaccines, is responsible for post-vaccination induced autoimmune diseases
11 in select patients. The risks have become so prolific that medical experts have coined a new umbrella
12 syndrome – Autoimmune/Inflammatory Syndrome Induced by Adjuvants ("ASIA") to refer to the
13 spectrum of immune-mediated diseases triggered by an adjuvant stimulus contained in vaccines, such
14 as aluminum. See e.g., YEHUDA SHOENFELD ET AL, EDS., VACCINES & AUTOIMMUNITY 2 (2015).

15 263. Indeed, even in animal studies, it has been revealed that aluminum adjuvants can induce
autoimmune disease in tested animals. By way of example, in a series of studies conducted by Lluís
Luján, DVM, Ph.D., and his colleagues, it was revealed that sheep injected with aluminum-containing
adjuvants commonly come down with severe autoimmune diseases and other adverse reactions.

19 264. Specific to the Gardasil vaccines, which contain adjuvants, including, amorphous
20 aluminum hydroxyphosphate sulfate (AAHS) and the previously undisclosed HPV L1 gene DNA
21 fragments, a number of mechanisms of action have been outlined (as discussed *infra*) as to how
22 Gardasil induces autoimmune disease in select patients.

23 265. Given the number of HPV strains that exist, a great part of the human population has
24 HPV, however, HPV by itself is generally not immunogenic, and generally does not evoke immune
25 responses. Indeed, HPV shares a high number of peptide sequences with human proteins, so that the
26 human immune system generally does not react against HPV in order to not harm self-proteins.
27 Immunotolerance thus generally blocks reactions against HPV in order to avoid autoimmune attacks
28 against the human proteins.

266. To induce anti-HPV immune reactions, Merck added various adjuvants, including
 amorphous aluminum hydroxyphosphate sulfate (AAHS), to the Gardasil vaccine. Adjuvants, such as
 aluminum, are inflammatory substances that hyperactivate the immune system. Adjuvants are thus
 the "secret sauce" used by Merck to hyperactivate the immune system and make HPV immunogenic.

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267. While adjuvants are added with the intent of destroying the HPV virus, they also can have the unintended result of rendering the immune system "blind" and unable to distinguish human proteins from HPV proteins – accordingly, human proteins that share peptide sequences with HPV are at risk of also being attacked by the vaccine.

268. While Gardasil causes immune hyperactivation and production of anti-HPV antibodies 9 to fend off certain strains of the HPV virus, it can also result in the immune system losing its ability to 10 differentiate human proteins from foreign proteins, causing the immune system to attack the body's 11 own proteins and organs. Because of the massive peptide commonality between HPV and human 12 proteins, the indiscriminate attack triggered by the Gardasil adjuvants will cause massive cross-13 reactions and dangerous attacks against human proteins, leading to a number of autoimmune diseases 14 manifested throughout the different organs of the body. This process is sometimes referred to as 15 "molecular mimicry." 16

269. In addition to "molecular mimicry," other mechanisms of action that explain how 17 Gardasil can induce autoimmune disease are "epitope spreading," whereby invading Gardasil 18 antigens, including the toxic aluminum adjuvant, accelerate autoimmune process by location 19 activation of antigen presenting cells and "bystander activation," wherein antigens and the aluminum 20 adjuvants in the Gardasil vaccine activate pre-primed autoreactive T cells, which can initiate 21 autoimmune disease (bystander activation of autoreactive immune T cells), or where virus-specific T 22 cells initiate bystander activation resulting in the immune system killing uninfected and unintended 23 neighboring cells. 24

25 270. Relevant to the injuries at issue in this case, when a person is lying down,
approximately one-quarter of their blood volume resides in the chest area. When the person stands
up, a significant amount of that blood shifts to the lower extremities. This causes impaired return of
blood flow to the heart which also reduces blood pressure. In healthy individuals, the autonomic

nervous system adjusts the heartrate to counteract this effect and the hemodynamic changes are 1 2 negligible. However, in individuals (such as Plaintiff) who are now suffering from dysautonomia or autonomic ailments, such as POTS or OI, the body's ability to adjust the heartrate and compensate for 3 the blood flow is corrupted resulting in a host of wide ranging symptoms, including but not limited to, 4 dizziness, lightheadedness, vertigo, woozy sensation, chronic headaches, vision issues due to the loss 5 of blood flow to the brain, light and sound sensitivity, loss of consciousness, shortness of breath, chest 6 pain, gastrointestinal issues, body pains, insomnia, and confusion and/or difficulty sleeping. In certain 7 cases of POTS, patients will also be diagnosed with other medical conditions, including but not 8 limited to, chronic fatigue syndrome and fibromyalgia. 9

271. Medical research has determined that certain dysautonomia diseases such as POTS and 10 OI have an autoimmune etiology. Norepinephrine, a key neurotransmitter of the sympathetic ("fight 11 or flight") system, exerts its mechanism of action by binding to receptors located in the smooth 12 muscle of the blood vessels and various organs, including the heart. These receptors include alpha-1, 13 alpha-2, beta-1, beta-2 and beta-3 receptors and, as a group, are generally known as the adrenergic 14 receptors. The adrenergic receptors, and other receptors, including but not limited to, the ganglionic 15 and muscarinic acetylcholine receptors are believed to be affected in certain cases of POTS and OI. 16 See e.g., Hongliang Li et al., Autoimmune Basis for Postural Tachycardia Syndrome, 3 J. AMERICAN 17 HEART ASSOC. e000755 (2014); Artur Fedorowski et al., Antiadrenergic Autoimmunity in Postural 18 Tachycardia Syndrome, 19 EUROPACE 1211 (2017); Mohammed Ruzieh et al., The Role of 19 Autoantibodies in the Syndromes of Orthostatic Intolerance: A Systematic Review, 51 SCANDINAVIAN 20 CARDIOVASCULAR J. 243 (2017); Shu-ichi Ikeda et al., Autoantibodies Against Autonomic Nerve 21 Receptors in Adolescent Japanese Girls after Immunization with Human Papillomavirus Vaccine, 2 22 ANNALS OF ARTHRITIS AND CLINICAL RHEUMATOLOGY 1014 (2019); William T. Gunning, Postural 23 Orthostatic Tachycardia Syndrome is Associated With Elevated G-Protein Coupled Receptor 24 Autoantibodies, 8 J. AMERICAN HEART ASSOC. e013602 (2019). 25

26 272. A variety of published medical journal articles have discussed the association between
 27 Gardasil and a myriad of serious injuries and have reported on patients developing POTS, OI,
 28 fibromyalgia and other symptoms of autonomic impairment following Gardasil vaccination. *See*

Svetlana Blitshetyn, Postural Tachycardia Syndrome After Vaccination with Gardasil, 17 EUROPEAN 1 2 J. OF NEUROLOGY e52 (2010); Svetlana Blitshetyn, Postural Tachycardia Syndrome Following Human Papillomavirus Vaccination, 21 EUROPEAN J. OF NEUROLOGY 135 (2014); Tomomi Kinoshita 3 et al., Peripheral Sympathetic Nerve Dysfunction in Adolescent Japanese Girls Following 4 Immunization With Human Papillomavirus Vaccine, 53 INTERNAL MEDICINE 2185 (2014); Louise S. 5 Brinth et al., Orthostatic Intolerance and Postural Tachycardia Syndrome As Suspected Adverse 6 Effects of Vaccination Against Human Papilloma Virus, 33 VACCINE 2602 (2015); Manuel Martinez-7 Lavin et al., HPV Vaccination Syndrome. A Questionnaire Based Study, 34 J. CLINICAL 8 RHEUMATOLOGY 1981 (2015); Louise S. Brinth et al., Is Chronic Fatigue Syndrome/Myalgic 9 Encephalomyelitis a Relevant Diagnosis in Patients with Suspected Side Effects to Human Papilloma 10 11 Virus Vaccine, 1 INT. J. OF VACCINE & VACCINATION 3 (2015); Jill R. Schofield et al., Autoimmunity, Autonomic Neuropathy, and HPV Vaccination, A Vulnerable Subpopulation, CLINICAL PEDIATRICS 12 (2017); Rebecca E. Chandler et al., Current Safety Concerns With Human Papillomavirus Vaccine: A 13 Cluster Analysis of Reports in VigiBase, 40 DRUG SAFETY 81 (2017); Svetlana Blitshetyn et al., 14 Autonomic Dysfunction and HPV Immunization An Overview, IMMUNOLOGIC RESEARCH (2018); and 15 Svetlana Blitshetyn, Human Papilloma Virus (HPV) Vaccine Safety Concerning POTS, CRPS and 16 *Related Conditions*, CLINICAL AUTONOMIC RESEARCH (2019). 17

18 273. In a 2017 review, Drs. Tom Jefferson and Lars Jørgensen criticized the European
19 Medicines Agency ("EMA") for turning a blind eye to the debilitating autoimmune injuries, including
20 CRPS and POTS that young women had suffered following vaccination with HPV vaccine. Tom
21 Jefferson et al., *Human Papillomavirus Vaccines, Complex Regional Pain Syndrome, Postural*22 Orthostatic Tachycardia Syndrome, and Autonomic Dysfunction – A Review of the Regulatory
23 Evidence from the European Medicines Agency, 3 INDIAN J. OF MED. ETHICS 30 (Jan. – March 2017).

24 274. In a separate article, the same authors describe their process for extracting data from not
25 only peer-reviewed journal publications, but also unpublished data from pharmaceutical company
26 clinical study reports and trial register entries from ClinicalTrials.gov, under the assumption that
27 "more than half of all studies are never published, and the published studies' intervention effects are
28 often exaggerated in comparison to the unpublished studies. This introduces reporting bias that

undermines the validity of systematic reviews. To address reporting bias in systematic reviews, it is 1 2 necessary to use industry and regulatory trial registers and trial data—in particular, the drug manufacturers' complete study programs." They found that 88 percent of industry studies were solely 3 industry funded and found serious deficiencies and variability in the availability of HPV vaccine study 4 data. For example, only half of the completed studies listed on ClinicalTrials.gov posted their results. 5 The clinical study reports the authors obtained confirmed that the amount of information and data are 6 vastly greater than that in journal publications. When the authors compared the data the EMA used 7 (which was provided by GlaxoSmithKline and Merck Sharp and Dohme) to conduct their review of 8 the relationship between HPV vaccination and both POTS and CRPS, the authors found that only 48 9 percent of the manufacturers' data were reported. According to the authors, "we find this very 10 disturbing." Lars Jørgensen et al., Index of the Human Papillomavirus (HPV) Vaccine Industry 11 Clinical Study Programmes and Non-Industry Funded Studies: A Necessary Basis to Address 12 *Reporting Bias in a Systematic Review*, 7 SYSTEMATIC REVIEW 8 (2018). 13

14 275. Likewise, in a recently released February 2020 peer-reviewed study, researchers who
15 analyzed the available clinical trial data for all HPV vaccines, which include the Gardasil vaccines and
16 another HPV vaccine currently only available in Europe, concluded that "HPV vaccines increased
17 serious nervous disorders." Lars Jørgensen et al., *Benefits and Harms of the Human Papillomavirus*18 (*HPV*) Vaccines: Systemic Review with Meta-Analyses of Trial Data from Clinical Study Reports, 9
19 SYSTEMATIC REVIEWS 43 (February 2020).

20 276. In addition, Jørgensen and his co-authors observed that, in reanalyzing the association
 21 between HPV vaccines and one specific autoimmune disease, POTS, the HPV vaccines were
 22 associated with a nearly two-fold increased risk of POTS. *Id.*

23 277. Jørgensen and his co-authors also noted many of the same shortcomings associated with
24 the Gardasil clinical trials as have already been discussed in this Complaint, including for example,
25 the fact that no true placebo was utilized by Merck as a comparator (i.e., the comparator/control used
26 by Merck in the Gardasil clinical trials contained aluminum adjuvant). The researchers noted that
27 "[t]he use of active comparators may have underestimated harms related to HPV vaccines," and that
28 "[t]he degree of harms might therefore be higher in clinical practice than in the trials." *Id*.

1 Jørgensen and his co-authors also noted that the clinical trials revealed that Gardasil 9 278. 2 induced more harms than Gardasil, which could be explained by the fact that Gardasil 9 contains more of the AAHS aluminum adjuvant (500 micrograms of AAHS in Gardasil-9 vs. 225 micrograms of 3 AAHS in Gardasil), and this dose-response relationship further corroborates the plausible claim that 4 the AAHS aluminum adjuvant is a culprit in causing adverse events. Id. 5

279. Other researchers, including Tomljenovic and Shaw, who have closely looked into 6 Gardasil, have opined that risks from the Gardasil vaccine seem to significantly outweigh the as yet 7 unproven long-term benefits. In their view, vaccination is unjustified if the vaccine carries any 8 substantial risk, let alone a risk of death, because healthy teenagers face an almost zero percent risk of 9 death from cervical cancer. 10

11

K. Merck has Concealed the Fact that Gardasil Increases the Risk of Fertility **Problems**

12 Merck has never tested the impact of the Gardasil vaccines on human fertility. 280. 13 281. Nevertheless, study volunteers reported devastating impacts on human fertility during 14 combined trials, offering substantial evidence that the vaccine may be causing widespread impacts on 15 human fertility, including increases in miscarriage, birth defects, premature ovarian failure and 16 premature menopause in girls and young women.

17 282. One of the serious adverse events now emerging in vaccinated girls, including teens, is 18 premature ovarian failure. See, e.g., D. T. Little and H. R. Ward, Adolescent Premature Ovarian 19 Insufficiency Following Human Papillomavirus Vaccination: A Case Series Seen in General Practice, 20 JOURNAL OF INVESTIGATIVE MEDICINE HIGH IMPACT, Case Reports 1-12 (Oct.-Dec. 2014); D. T. Little 21 and H. R. Ward, Premature ovarian failure 3 years after menarche in a 16-year-old girl following 22 human papillomavirus vaccination, BMJ CASE REPORTS (September 30, 2012).

23

283. Premature ovarian failure can occur after aluminum destroys the maturation process of 24 the eggs in the ovaries.

25 284. Fertility has plummeted among American women following the 2006 mass introduction 26 of the Gardasil vaccine. This is most evident in teen pregnancy statistics where numbers have more 27 than halved since 2007.

28

285. The total fertility rate for the United States in 2017 continued to dip below what is
 needed for the population to replace itself, according to a report by the National Center of Health
 Statistics issued in January 2019, and the rate for women 15 to 44 fell another 2 percent between 2017
 and 2018.

5

L. There were an Increased Number of Deaths in the Gardasil Studies

6 286. Merck's own preliminary studies predicted that Gardasil would kill and injure far more
7 Americans than the HPV virus, prior to the introduction of the vaccine.

8 287. The average death rate in young women in the U.S. general population is 4.37 per
9 10,000. See Brady E. Hamilton et al., "Births: Provisional Data for 2016," *Vital Statistics Rapid*10 *Release, Report No. 002*, June 2017.

288. The Gardasil pooled group had a death rate of 8.5 per 10,000, or almost double the
background rate in the U.S.

13 Age adjusted Incidence of death per 10,000 girls age 15-26 14 8.5 9 15 8 7.2 7 16 6 CDC 4.37 5 17 Cervical cance 4 leath rate 0.23 per 10,000 3 18 2 1 19 0 A11 C Gardasil AAHS Co Background CDC rate 4.37 source: National Vital Statistics Report Vol. 53 2002 page 24.37 Gardasil rate 8.5: 10/11.778. AAHS control rate 7.2: 7/9.68038 20 Cervical cancer mortality: 2.3 per 100,000 spurce: National Cancer Institute SEER Cancer Statistics Review 201539 21 22 289. When Merck added in deaths from belated clinical trials, the death rate jumped to 13.3 per 10,000 (21 deaths out of 15,706). 23 Merck dismissed all deaths as coincidences. 290. 24 25 291. The total number of deaths was 21 in the HPV vaccine group and 19 in the comparator 26 (AAHS) groups. The death rate among vaccine recipients was 13.3 per 10,000, or 133 per 100,000 27 292. (21/15,706).28 45 COMPLAINT

293. To put this in perspective, the death rate from cervical cancer in the United States is 2.3
 per 100,000 women. This means that, according to Merck's own data, a girl is 58 times more likely to
 die from Gardasil than from cervical cancer.

4

M. Post-Marketing Injuries -- The Raft of Injuries Seen in Merck's Clinical Trials Has Now Become A Population-Wide Chronic Disease Epidemic

5 294. By 2010, reports coming in from all over the world linked the Gardasil vaccine to
6 bizarre and troubling symptoms.

7

295. Many Gardasil survivors will have lifelong handicaps.

8 296. The severe adverse events from the Gardasil vaccination, seen since its widespread
 9 distribution, are similar to those injuries that Merck covered up during its clinical trials. They include
 10 autoimmune diseases, suicides, deaths, premature ovarian failures, reproductive problems, infertility,
 11 cervical cancer, sudden collapse, seizures, multiple sclerosis, strokes, heart palpitations, chronic
 12 muscle pain, complex regional pain syndrome, and weakness.

13 297. Other frequently reported injuries include disturbances of consciousness; systemic pain 14 including headache, myalgia, arthralgia, back pain and other pain; motor dysfunction, such as 15 paralysis, muscular weightiness, and involuntary movements; numbness, and sensory disturbances; 16 autonomic symptoms including hypotension, tachycardia, nausea, vomiting, and diarrhea; respiratory 17 dysfunction, including dyspnea, and asthma; endocrine disorders, such as menstrual disorder and 18 hypermenorrhea; and lastly, hypersensitivity to light, heart palpitations, migraine headaches, 19 dizziness, cognitive deficits, personality changes, vision loss, joint aches, headaches, brain 20 inflammation, chronic fatigue, death, and severe juvenile rheumatoid arthritis.

21 298. The data show that Gardasil is yielding far more reports of adverse events than any
22 other vaccine. For example, Gardasil had 8.5 times more emergency room visits, 12.5 times more
23 hospitalizations, 10 times more life-threatening events, and 26.5 more disabilities than Menactra,
24 another vaccine with an extremely high-risk profile.

25 299. As of December 2019, there have been more than 64,000 Gardasil adverse events
26 reported to the FDA's Vaccine Adverse Event Reporting System ("VAERS") since 2006.

27 28

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300. Moreover, studies have shown that only approximately 1 percent of adverse events are
 actually reported to FDA's voluntary reporting systems, thus, the true number of Gardasil adverse
 events in the United States may be as high as 6.4 million incidents.

301. The Vaccine Injury Compensation Program has paid out millions of dollars in damages
for Gardasil-induced injuries and deaths.

302. The adverse events also include deaths. Parents, doctors, and scientists have reported
hundreds of deaths from the Gardasil vaccine, post-marketing.

8 303. In order to conceal Gardasil's link to the deaths of teenagers, Merck has submitted
9 fraudulent reports to VAERS, and posts fraudulent and misleading statements on its Worldwide
10 Adverse Experience System.

304. For example, Merck attributed the death of a young woman from Maryland, Christina
Tarsell, to a viral infection. Following years of litigation, a court determined that Gardasil caused
Christina's death. There was no evidence of viral infection. Merck invented this story to deceive the
public about Gardasil's safety.

305. Merck submitted fraudulent information about Christina Tarsell's death to its
Worldwide Adverse Experience System and lied to the FDA through the VAERS system. Merck
claimed that Christina's gynecologist had told the company that her death was due to viral infection.
Christina's gynecologist denied that she had ever given this information to Merck. To this day, Merck
has refused to change its false entry on its own reporting system.

20

N. The Gardasil Vaccines' Harms Are Not Limited to the United States, Rather the Vaccines Have Injured Patients All Over the World

In Light of Gardasil's Serious and Debilitating Adverse Events, the

Japanese Government Rescinded Its Recommendation that Girls

306. Gardasil is used widely in the international market. Widespread global experience has
likewise confirmed that the vaccine causes serious adverse events with minimal proven benefit.

- 23307. According to the World Health Organization's Adverse Event Databases, there have
- 24 been more than 100,000 serious adverse events associated with Gardasil, outside the Americas. See
- 25 WHO Vigibase database, keyword Gardasil: http://www.vigiaccess.org.

Receive Gardasil

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308. In Japan, a country with a robust history of relative honesty about vaccine side effects,
 the cascade of Gardasil injuries became a public scandal.

3 309. Japan's health ministry discovered adverse events reported after Gardasil were many
times higher than other vaccines on the recommended schedule. These included seizures, severe
headaches, partial paralysis, and complex regional pain syndrome. See Hirokuni Beppu et al., *Lessons Learnt in Japan From Adverse Reactions to the HPV Vaccine: A Medical Ethics Perspective*, 2
INDIAN J MED ETHICS 82 (April-June 2017).

310. Japanese researchers found that the adverse events rate of the HPV vaccine was as high
as 9 percent, and that pregnant women injected with the vaccine aborted or miscarried 30 percent of
their babies. *See* Ministry of Health, Labour and Welfare, Transcript "The Public Hearing on Adverse
Events following HPV vaccine in Japan," February 26, 2014.

12 311. The injuries caused the Japanese government to rescind its recommendation that girls
13 receive the HPV vaccine.

312. Japan withdrew its recommendation for Gardasil three months after it had added the
vaccine to the immunization schedule, due to "an undeniable causal relationship between persistent
pain and the vaccination."

313. Uptake rates for the vaccine in Japan are now under 1 percent, compared to 53.7 percent
fully vaccinated teenaged girls in the United States.

19 314. In late 2016 Japanese industry watchdog, MedWatcher Japan issued a scathing letter
20 faulting the WHO for failing to acknowledge the growing body of scientific evidence demonstrating
21 high risk of devastating side effects.

315. In 2015, the Japanese Association of Medical Sciences issued official guidelines for
 managing Gardasil injuries post-vaccination.

316. That same year, the Japanese Health Ministry published a list of medical institutions
 where staffs were especially trained to treat patients who had sustained Gardasil-induced injuries.

317. The Japanese government also launched a series of special clinics to evaluate and treat
 illnesses caused by the Gardasil vaccines.

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Case 2:22-cv-00160 Document 1 Filed 04/01/22 Page 49 of 83 PageID #: 49 1 The president of the Japanese Association of Medical Sciences stated that there was no 318. 2 proof that the vaccines prevent cancer. These were developments that Merck was extremely anxious to suppress. 3 319. Merck hired the think tank, the Center for Strategic and International Studies ("CSIS") 320. 4 and Professor Heidi Larson of the Vaccine Confidence Project in London, to assess the reasons for the 5 Japanese situation. The overall conclusion was that the symptoms the girls were suffering from were 6 psychogenic in nature and were a result of rumors spread online. In essence, Merck blamed the 7 victims for the Gardasil-induced adverse events in Japan. 8 ii. **Denmark Has Opened Specialized Clinics Specifically Focused on** 9 Treating Gardasil-Induced Injuries, Including Gardasil-Induced **Autoimmune Diseases** 10 321. In March 2015, Denmark announced the opening of five new "HPV clinics" to treat 11 children injured by Gardasil vaccines. Over 1,300 cases flooded the HPV clinics shortly after 12 opening. See Zosia Chustecka, Chronic Symptoms After HPV Vaccination: Danes Start Study, 13 MEDSCAPE (November 13, 2015). 14 iii. Gardasil-Induced Adverse Events Caused the Government in 15 Colombia to Conclude that Gardasil Would No Longer Be Mandatory 16 322. In Colombia, more than 800 girls in the town of El Carmen de Bolivar reported 17 reactions ranging from fainting to dizziness to paralysis in March of 2014, following vaccination with 18 Gardasil. 19 323. With protests erupting across the country, the Colombian attorney general asked the 20 Constitutional Court to rule on a lower court ruling on the outcome of a case of an injured girl. 21 324. In 2017, in response to an unresolved case, Colombia's constitutional court, ruled that 22 the Colombian government could not infringe on the bodily integrity of its citizens. This decision 23 meant that the government could not require the HPV vaccine to be mandatory. 24 India Halted Gardasil Trials and Accused Merck of Corruption iv. 25 After the Death of Several Young Girls Who were Participants in the Trial 26 325. Seven girls died in the Gardasil trials in India coordinated by Merck and the Gates 27 Foundation. A report by the Indian Parliament accused the Gates Foundation and Merck of 28 49

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conducting "a well-planned scheme to commercially exploit" the nation's poverty and powerlessness
 and lack of education in rural India in order to push Gardasil. See 72nd Report on the Alleged
 Irregularities in the Conduct of Studies Using Human Papilloma Virus (HPV) Vaccine by Programme
 for Appropriate Technology in Health (PATH) in India (August 2013).

326. The report alleges that Merck (through PATH, to whom it supplied vaccines) and the
Gates Foundation resorted to subterfuge that jeopardized the health and well-being of thousands of
vulnerable Indian children. The parliamentary report makes clear that the clinical trials could not have
occurred without Merck corrupting India's leading health organizations. *Id*.

327. The Report accused PATH, which was in collaboration with Merck, of lying to illiterate
tribal girls to obtain informed consent, widespread forging of consent forms by Merck operatives,
offering financial inducements to participate, and providing grossly inadequate information about
potential risks. *Id.*

328. Many of the participants suffered adverse events including loss of menstrual cycles and
psychological changes like depression and anxiety. According to the report: PATH's "sole aim has to
been to promote the commercial interests of HPV vaccine manufacturers, who would have reaped a
windfall of profits had they been successful in getting the HPV vaccine included in the universal
immunization program of the country... This [conduct] is a clear-cut violation of the human rights of
these girls and adolescents." *Id.*

329. A 2013 article in the *South Asian Journal of Cancer* concludes that the HPV vaccine
program is unjustifiable. "It would be far more productive to understand and strengthen the reasons
behind the trend of decreasing cervical cancer rates than to expose an entire population to an uncertain
intervention that has not been proven to prevent a single cervical cancer or cervical cancer death to
date." *See* Sudeep Gupta, *Is Human Papillomavirus Vaccination Likely to be a Useful Strategy in India?* 2 SOUTH ASIAN J CANCER 194 (October-December 2014).

330. The article goes on to say: "A healthy 16-year-old is at zero immediate risk of dying
from cervical cancer, but is faced with a small, but real risk of death or serious disability from a
vaccine that has yet to prevent a single case of cervical cancer... There is a genuine cause for concern
regarding mass vaccination in this country." *Id.*

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1	331. In April 2017, the Indian government blocked the Gates Foundation from further					
2	funding of the Public Health Foundation of India and other non-governmental organizations,					
3	effectively barring them from influencing India's national vaccine program. See Nida Najar, India's					
4	Ban on Foreign Money for Health Group Hits Gates Foundation, THE NEW YORK TIMES, April 20,					
5	2017.					
6		O. Merck's Fraud Has Paid Off Handsomely Resulting in Over \$3 Billion in Gardasil Sales Annually				
7	332.	Merck's corruption and fraud in researching, testing, labeling, and promoting Gardasil				
8	have paid off handsomely.					
9	333.	Presently, two doses of Gardasil 9 typically cost about \$450, plus the cost of two office				
10	visits.					
11	334.	By comparison, the cost of the DTaP vaccine is about \$25 per dose.				
12	2 335. The HPV vaccine is the most expensive vaccine on the market.					
13	336. Since approximately 1 in 42,000 American women die of cervical cancer annually, the					
14	cost of avoiding a single death is over \$18 million, assuming the Gardasil vaccine is 100 percent					
15	effective.					
16	337.	In 2018, the Gardasil vaccines made \$2.2 billion for Merck in the U.S. alone.				
17	338.	In 2019, Merck made \$3.7 billion in worldwide revenues from the Gardasil vaccines.				
18	339.	Gardasil is Merck's most lucrative vaccine and its third-highest selling product.				
19	340.	Gardasil is crucial to Merck's overall financial health. Merck identifies Gardasil as one				
20	of its "key products," meaning that any change in Gardasil's cash flow affects the corporation as a					
21	whole.					
22	341.	Merck's 10-K financial reports note that, for example, the discovery of a previously				
23	unknown side effect, or the removal of Gardasil from the market, would hurt Merck's bottom line.					
24	III.	I.L. Sustained Autoimmune Disease, Autonomic Dysfunction and Other Serious Injuries, as A Result of Her Gardasil Injection				
25		A. Gardasil and Its Ingredients Caused Plaintiff's Autoimmune Disease and Other				
26		Related Injuries and Has Resulted in Her Suffering from Severe, Debilitating, Disabling and Painful Chronic Injuries				
27	342.	Plaintiff was eleven (11) years old when she received her first dosage of Gardasil on				
28	September 12	, 2019.				
		51 COMPLAINT				

343. Plaintiff's mother, Elizabeth Landers, agreed to her daughter receiving the Gardasil
 injection after having been exposed to marketing by Merck, that Gardasil is very safe, that Gardasil
 prevents cancer and that teenagers must get the Gardasil vaccine. Plaintiff's mother relied upon
 Merck's ubiquitous representations concerning the safety and efficacy of the Gardasil vaccine, in
 consenting to her daughter's Gardasil vaccination.

344. Prior to receiving her Gardasil injection, Plaintiff had no autoimmune diseases, and no
autonomic issues. I.L. was attending school, playing softball, cheering through Upwards Sports
Program, and playing golf for her middle school team. She loved going to school, walking her dog,
playing sports and going to church.

345. Before receiving this vaccination, I.L.'s medical history notes asthma, positive ANA,
corrective eye surgery, and amplified musculoskeletal pain, however she was still active and carrying
out her normal activities. Indeed, the encounter record done immediately before vaccination states,
"Encounter for routine child health examination without abnormal findings."

346. On September 12, 2019, Plaintiff's health care provider in Parkersburg, West Virginia
recommended that Plaintiff receive the Gardasil vaccine, which was stated as a safe and effective
vaccine for preventing cervical cancer. In light of the doctor's recommendations, as well as Merck's
relentless marketing and advertising messages, to which Plaintiff's mother had been exposed
concerning the safety and efficacy of Gardasil, Plaintiff's mother consented to Plaintiff being injected
with the "cervical cancer vaccine," Gardasil.

347. On or about October 3, 2019, I.L.'s school nurse called Ms. Landers to inform her that
I.L's heart rate was 160 and her blood pressure was low and that she should be taken to an ER for
assessment. From WVU emergency room, I.L. went by ambulance to Nationwide Children's Hospital
in Columbus, Ohio for stroke and decreasing neurological activity. Her body went into septic shock.

348. After a four (4) day and three (3) night hospitalization, I.L. was given papers on
Postural Orthostatic Tachycardic Syndrome (POTS) and discharged home. I.L. required the use a
walker to help her move. Plaintiff did not have the ability to move like she did before, could not get
up alone, and could barely walk without fainting or collapsing.

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1 After being released from the hospital, I.L. attempted to return to school but was instead 349. placed on a homebound schedule. 2

350. On October 16, 2019, I.L. was again taken to the hospital, suffering from palpitations, 3 dizziness and headaches, similar to her previous POTS episodes. 4

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351. On November 8, 2019, I.L. had a POTS flare during her sister's birthday party. It led to seizures. She was carried out of the skating rink and taken to the local ER. They completed blood 6 work and sent I.L. to Nationwide Children's Hospital by ambulance. I.L's seizures lasted 14 hours on 7 and off throughout the night. She went home the next morning. 8

352. On November 13, 2019, I.L. attended a Wednesday night church service. During the 9 service, I.L. started having seizures. I.L. was transported to Marietta Memorial in Belpre, Ohio, tested, 10 and referred once again sent to Nationwide Children's Hospital. This time her seizures lasted 12 hours 11 on and off throughout the night. She was monitored for neurological activity and was released again 12 the following morning. 13

353. As the months progressed, so did Plaintiff's injuries. She was seen by multiple 14 physicians and specialists for her complaints which now included: fainting spells, seizures, 15 hypotension, sensitivity to lights and sounds, sensitivity to hot and cold, dizziness, brain fog, nausea, 16 and unprovoked triggering of the "fight or flight" response. 17

354. As a result of her post-Gardasil symptoms, Plaintiff was unable to engage in normal 18 activities that a normal young person would enjoy. I.L was unable to return to school and was placed 19 on a homebound schedule through 2019. In January 2020, Plaintiff attempted to return to school, but 20 her condition did not allow her to attend on a regular basis. Plaintiff has further been forced to forgo 21 the extracurricular activities she enjoyed before her Gardasil vaccination, including academic 22 competitions, softball, cheering, golf and other sports. Because of her condition Plaintiff is unable to 23 independently care for her basic needs, and the severity of her symptoms have left her unable to 24 shower. 25

355. Based upon her chronic and severe post-Gardasil symptoms, Plaintiff has been 26 diagnosed with various medical conditions, including but not limited to, Postural Orthostatic 27

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Tachycardia Syndrome, progressively worsening vision, seizures, and other autonomic nervous
 system dysfunctions.

3 356. As previously discussed, the medical literature has documented other patients who, like Plaintiff, have suffered serious autonomic dysfunctions, and who experienced the same side effects as 4 5 those Plaintiff has suffered, and who were diagnosed with Gardasil-induced autonomic diseases. See E. Israeli et al., Adjuvants and Autoimmunity, 18 LUPUS 1217 (2009); Darja Kanduc, Quantifying the 6 7 Possible Cross-Reactivity Risk of an HPV16 Vaccine, 8 JOURNAL OF EXPERIMENTAL THERAPEUTICS AND ONCOLOGY 65 (2009); Svetlana Blitshetyn, Postural Tachycardia Syndrome After Vaccination 8 9 with Gardasil, 17 EUROPEAN J. OF NEUROLOGY e52 (2010); Darja Kanduc, Potential Cross-Reactivity Between HPV16 L1 Protein and Sudden Death Associated Antigens, 9 JOURNAL OF EXPERIMENTAL 10 11 THERAPEUTICS AND ONCOLOGY 159 (2011); Deirdre Little et al., Premature ovarian failure 3 years 12 after menarche in a 16-year-old girl following human papillomavirus vaccination, BRIT. MED. J. CASE 13 REPORTS (2012); Serena Colafrancesco et al., Human Papilloma Virus Vaccine and Primary Ovarian 14 Failure: Another Facet of the Autoimmune Inflammatory Syndrome Induced by Adjuvants, 70 AM. J. REPRODUCTIVE IMMUNOLOGY 309 (2013); Maurizo Rinaldi et al., Anti-Saccharomyces Cerevisiae 15 16 Autoantibodies in Autoimmune Diseases: from Bread Baking to Autoimmunity, 45 CLINICAL REVIEWS 17 IN ALLERGY AND IMMUNOLOGY 152 (October 2013); Svetlana Blitshetyn, Postural Tachycardia Syndrome Following Human Papillomavirus Vaccination, 21 EUROPEAN J. OF NEUROLOGY 135 18 19 (2014); Tomomi Kinoshita et al., Peripheral Sympathetic Nerve Dysfunction in Adolescent Japanese 20 Girls Following Immunization With Human Papillomavirus Vaccine, 53 INTERNAL MEDICINE 2185 21 (2014); Christopher A. Shaw et al., *Aluminum-Induced Entropy in Biological Systems: Implications* 22 for Neurological Disease, JOURNAL OF TOXICOLOGY (2014); Louise S. Brinth et al., Orthostatic 23 Intolerance and Postural Tachycardia Syndrome As Suspected Adverse Effects of Vaccination Against 24 Human Papilloma Virus, 33 VACCINE 2602 (2015); Manuel Martinez-Lavin et al., HPV Vaccination 25 Syndrome. A Questionnaire Based Study, 34 J. CLINICAL RHEUMATOLOGY 1981 (2015); Louise S. Brinth et al., Is Chronic Fatigue Syndrome/Myalgic Encephalomyelitis a Relevant Diagnosis in 26 27 Patients with Suspected Side Effects to Human Papilloma Virus Vaccine, 1 INT. J. OF VACCINE & 28 VACCINATION 3 (2015); Jill R. Schofield et al., Autoimmunity, Autonomic Neuropathy, and HPV

Vaccination, A Vulnerable Subpopulation, CLINICAL PEDIATRICS (2017); Rebecca E. Chandler et al., 1 2 Current Safety Concerns With Human Papillomavirus Vaccine: A Cluster Analysis of Reports in VigiBase, 40 DRUG SAFETY 81 (2017); Svetlana Blitshetyn et al., Autonomic Dysfunction and HPV 3 Immunization An Overview, IMMUNOLOGIC RESEARCH (2018); and Svetlana Blitshetyn, Human 4 5 Papilloma Virus (HPV) Vaccine Safety Concerning POTS, CRPS and Related Conditions, CLINICAL AUTONOMIC RESEARCH (2019); Lars Jørgensen et al., Benefits and Harms of the Human 6 7 Papillomavirus (HPV) Vaccines: Systemic Review with Meta-Analyses of Trial Data from Clinical 8 Study Reports, 9 SYSTEMATIC REVIEWS 43 (February 2020). 9 357. Plaintiff contends that her Gardasil injection caused her to develop serious and debilitating injuries, including but not limited to autonomic, neurological, heterogenous autoimmune 10 11 disease, POTS, and dysautonomia, as well as a constellation of adverse symptoms, complications, 12 injuries, and other adverse events, many of which are alleged herein and all of which were caused by Gardasil or otherwise linked to her Gardasil-induced autoimmune disorder. 13 B. "It is Not Revolutions and Upheavals That Clear the Road to New and Better 14 Days, But Revelations, Lavishness and Torments of Someone's Soul, Inspired 15 and Ablaze." - Boris Pasternak, After the Storm 358. Pursuant to Section 300aa-11(a) of the National Vaccine Injury Compensation 16 Program: "No person may bring a civil action for damages against a vaccine administrator or 17 manufacturer in a State or Federal court for damages arising from a vaccine-related injury ... 18 associated with the administration of a vaccine unless a petition has been filed, in accordance 19 with section 300aa-16 of this title, for compensation under the Program for such injury ... and (I) the 20 United Stated Court of Federal Claims has issued a judgment under section 300aa-12 of this title on 21 22 such petition and (II) such person elects under section 300aa-21(a) to file such an action." See 42 U.S.C. §§ 300aa–11(a)(2)(A). 23 359. Title 42, Section 300aa-16 (c) further states: "If a petition is filed under section 300aa-24 11 of this title for a vaccine-related injury or death, limitations of actions under State law shall be 25 stayed with respect to a civil action brought for such injury or death for the period beginning on the 26 date the Petition is filed and ending on the date...an election is made under section 300aa-21(a) of this 27

> 55 COMPLAINT

title to file the civil action ..." See 42 U.S.C. §§ 300aa–16(c).

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360. In full compliance with the aforementioned federal law, Plaintiff duly filed her petition
 with the U.S. Court of Federal Claims seeking compensation for her Gardasil vaccine-related injuries
 under the National Vaccine Injury Compensation Program. The Order Concluding Proceedings was
 filed on February 2, 2022.

361. Having complied with National Vaccine Injury Compensation Program administrative
procedure and having duly filed her election to proceed with a civil action, Plaintiff hereby timely
initiates the instant action against Merck, the manufacturer and promoter of the Gardasil vaccines
which caused her debilitating injuries. Through this civil action, Plaintiff seeks to hold Merck
accountable for its negligent, reckless, and fraudulent conduct and she seeks full compensation from
Merck for the physical and emotional injuries and harms she sustained as a result of Gardasil.

COUNT ONE

NEGLIGENCE

CAUSES OF ACTION

14 362. Plaintiff incorporates by reference all other paragraphs of this Complaint as if fully set15 forth herein and further alleges:

363. Merck is the researcher, manufacturer, labeler, and promoter of the Gardasiland the subsequent Gardasil 9 vaccines.

18 364. Merck marketed Gardasil to patients, including teenagers such as Plaintiff and her
19 medical providers.

365. Merck had a duty to exercise reasonable care in the research, manufacture, marketing,
advertisement, supply, promotion, packaging, sale, and distribution of Gardasil, including the duty to
take all reasonable steps necessary to research, manufacture, label, promote and/or sell a product that
was not unreasonably dangerous to consumers, users, and other persons coming into contact with the
product.

366. At all times relevant to this litigation, Merck had a duty to exercise reasonable care in
 the marketing, advertising, and sale of Gardasil. Merck's duty of care owed to consumers and the
 general public included providing accurate, true, and correct information concerning the efficacy and

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risks of Gardasil and appropriate, complete, and accurate warnings concerning the potential adverse
 effects of Gardasil and its various ingredients and adjuvants.

3 367. At all times relevant to this litigation, Merck knew or, in the exercise of reasonable care,
4 should have known of the hazards and dangers of Gardasil and specifically, the serious, debilitating
5 and potentially fatal adverse events associated with Gardasil, including but not limited to autoimmune
6 diseases (including, but not limited to, POTS and OT), fibromyalgia, increased risk of cancer
7 (including cervical cancer, which was the very cancer it was promoted as preventing), and death.

8 368. Accordingly, at all times relevant to this litigation, Merck knew or, in the exercise of
9 reasonable care, should have known that use of Gardasil could cause Plaintiff's injuries and thus
10 created a dangerous and unreasonable risk of injury to the users of these products, including Plaintiff.

369. Merck knew or, in the exercise of reasonable care, should have known that its
negligently and poorly performed clinical trials and studies were insufficient to test the true long-term
safety and efficacy of Gardasil.

370. Merck also knew, or, in the exercise of reasonable care, should have known that its
targeted consumers and patients (who were pre-teen and teen children), the parents of these patients
and the children's medical providers were unaware of the true risks and the magnitude of the risks
associated with Gardasil and the disclosed and undisclosed ingredients of Gardasil.

371. As such, Merck breached its duty of reasonable care and failed to exercise ordinary care
in the research, development, manufacturing, testing, marketing, supply, promotion, advertisement,
packaging, labeling, sale, and distribution of Gardasil, in that Merck manufactured and produced a
defective and ineffective vaccine, knew or had reason to know of the defects and inefficacies inherent
in its products, knew or had reason to know that a patient's exposure to Gardasil created a significant
risk of harm and unreasonably dangerous side effects, and failed to prevent or adequately warn of
these defects, risks and injuries.

372. Merck failed to appropriately and adequately test the safety and efficacy of Gardasil and
 its individual ingredients and adjuvants.

27 373. Despite the ability and means to investigate, study, and test its products and to provide
 28 adequate warnings, Merck has failed to do so. Indeed, Merck has wrongfully concealed information

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and has further made false and/or misleading statements concerning the safety and efficacy of 1 2 Gardasil.

3	374.	374. Merck's negligence is outlined in detail in this Complaint and included, among					
4	other things:						
5	a)	Manufacturing, producing, promoting, creating, researching, labeling, selling,					
6		and/or distributing Gardasil without thorough and adequate pre-and post-market					
7		testing and studies;					
8	b)	Manufacturing, producing, promoting, researching, labeling, selling, and/or					
9		distributing Gardasil while negligently and intentionally concealing and failing					
10		to accurately and adequately disclose the results of the trials, tests, and studies of					
11		Gardasil, and, consequently, the lack of efficacy and risk of serious harm					
12		associated with Gardasil;					
13	c)	Failing to undertake sufficient studies and conduct necessary tests to determine					
14		the safety of the ingredients and/or adjuvants contained within Gardasil, and the					
15		propensity of these ingredients to render Gardasil toxic, increase the toxicity of					
16		Gardasil, whether these ingredients are carcinogenic or associated with					
17		autoimmune diseases and other injures;					
18	d)	Negligently designing and conducting its clinical trials so as to prevent the					
19		clinical trials from revealing the true risks, including but not limited to, long					
20		terms risks and risks of autoimmune diseases associated with Gardasil;					
21	e)	Negligently designing and conducting its clinical trials so as to mask the true					
22		risks, including but not limited to, long terms risks and risks of autoimmune					
23		diseases and cancers associated with Gardasil;					
24	f)	Failing to test Gardasil against a true inert placebo and lying to the public that					
25		Gardasil was tested against a placebo, when in reality, all, or nearly all, studies					
26		used a toxic placebo that included the aluminum adjuvant AAHS;					
27	g)	Failing to have a sufficient number of studies for the targeted patient population					
28		which included pre-teen girls (and boys) between the ages of nine and 12;					
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1	h)	Not using the commercial dosage (and instead using a lower dosage of the
2		adjuvant and ingredients) in one of the key clinical trials used to obtain licensing
3		for the commercial dosage of Gardasil;
4	i)	Using restrictive exclusionary criteria in the clinical study patient population
5		(including for example, the exclusion of anyone who had prior abnormal Pap
6		tests, who had a history of immunological or nervous system disorders, or was
7		allergic to aluminum or other ingredients), but then not revealing or warning
8		about these exclusionary criteria in the label and knowing that, for most of these
9		ingredients and allergies, there are limited resources for the public to test for
10		such allergies in advance of being vaccinated;
11	j)	Negligently designing and conducting its trials so as to create the illusion of
12		efficacy when in reality the Gardasil Vaccines have not been shown to be
13		effective against preventing cervical and anal cancer;
14	k)	Failing to use reasonable and prudent care in the research, manufacture, labeling
15		and development of Gardasil so as to avoid the risk of serious harm associated
16		with the prevalent use of Gardasil;
17	1)	Failing to provide adequate instructions, guidelines, warnings, and safety
18		precautions to those persons who Merck could reasonably foresee would use
19		and/or be exposed to Gardasil;
20	m)	Failing to disclose to Plaintiff and her medical providers and to the general
21		public that Gardasil is ineffective when used in patients who have previously
22		been exposed to HPV, and also failing to disclose that Gardasil
23		actually increases the risk of cervical cancer, including in any child or patient
24		who has previously been exposed to HPV;
25	n)	Failing to disclose to Plaintiff and her medical providers and to the general
26		public that use of and exposure to Gardasil presents severe risks of cancer
27		(including cervical cancer, the very cancer it is promoted as preventing), fertility
28		problems, autoimmune diseases and other grave illnesses as alleged herein;
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o)	Failing to disclose to Plaintiff and her medical providers and to the general
	public that use of and exposure to Gardasil presents severe risks of triggering
	and increasing the risk of various autoimmune diseases, including but not
	limited to POTS and OI;

p) Failing to disclose to Plaintiff and her medical providers and to the general public that, contrary to Merck's promotion of the vaccine, Gardasil has not been shown to be effective at preventing cervical cancer and that the safest and most effective means of monitoring and combating cervical cancer is regular testing, including Pap tests;

q) Representing that Gardasil was safe and effective for its intended use when, in fact, Merck knew or should have known the vaccine was not safe and not effective for its intended use;

 r) Falsely advertising, marketing, and recommending the use of Gardasil, while concealing and failing to disclose or warn of the dangers Merck knew to be associated with or caused by the use of Gardasil;

s) Falsely promoting Gardasil as preventing cervical cancer when Merck knows that it has not done any studies to demonstrate that Gardasil prevents cervical cancer and, indeed, its clinical studies revealed that Gardasil actually increases the risk of cervical cancer;

t) Engaging in false advertising and disease mongering by scaring parents and children into believing that cervical and anal cancer is far more prevalent than it really is; that all cervical and anal cancer was linked to HPV; that Gardasil prevented cervical and anal cancer, when in reality none of these representations were true as cervical cancer rates were declining in the United States due to Pap testing and Gardasil has not been shown to prevent against all strains of HPV that are associated with cervical and anal cancer;

Failing to disclose all of the ingredients in Gardasil, including but not limited to 1 u) 2 the fact that Gardasil contains dangerous HPV L1-DNA fragments and that these DNA fragments could act as a Toll-Like Receptor 9 (TLR9) agonist -3 further adjuvanting the vaccine and making it more potent and dangerous; 4 v) Declining to make any changes to Gardasil's labeling or other promotional 5 materials that would alert consumers and the general public of the true risks and 6 defects of Gardasil; 7 Systemically suppressing or downplaying contrary evidence about the risks, 8 w) incidence, and prevalence of the side effects of the Gardasil Vaccines by, inter 9 alia, orchestrating the retraction of peer-reviewed and published studies and 10 vilifying and attempting to ruin the careers of any scientists who openly question 11 Gardasil's safety and efficacy. 12 375. Merck knew and/or should have known that it was foreseeable that patients, such as 13 Plaintiff, would suffer injuries as a result of Merck's failure to exercise ordinary care in the 14 manufacturing, marketing, labeling, distribution, and sale of Gardasil. 15 Plaintiff and, upon information and belief, her medical providers, did not know the true 376. 16 nature and extent of the injuries that could result from the intended use of and/or exposure to Gardasil 17 or its adjuvants and ingredients. 18 377. Merck's negligence was the proximate cause of the injuries, harm, and economic losses 19 that Plaintiff suffered, and will continue to suffer, as described herein. 20 378. Had Merck not engaged in the negligent and fraudulent conduct alleged herein and/or 21 had Merck via its labeling, advertisements, and promotions provided adequate and truthful warnings 22 and properly disclosed and disseminated the true risks, limitations, and lack of efficacy associated 23 with Gardasil to medical providers, patients and the public, then upon information and belief, 24

25 Plaintiff's medical providers would not have offered or recommended Gardasil to Plaintiff.

26 Moreover, even if after Merck's dissemination of truthful information concerning the true risks and

27 efficacy limitation of Gardasil, Plaintiff's medical providers had offered Gardasil, then upon

28 information and belief, the providers would have heeded any warnings issued by Merck and relayed

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to Plaintiff the safety risks and efficacy limitations that Merck should have warned her about, but
 failed to do so. Had Plaintiff been informed of the true risks and efficacy limitation concerning
 Gardasil, either through her medical providers or through Merck's ubiquitous direct-to-consumer
 promotional marketing, on which Plaintiff relied, then Plaintiff would never have consented to
 Plaintiff being injected with Gardasil.

6 379. As a proximate result of Merck's wrongful acts and omissions and its negligent and
7 fraudulent testing, labeling, manufacturing, marketing and promotion of Gardasil, Plaintiff has
8 suffered and continues to suffer severe and permanent physical injuries, and associated symptomology
9 and has suffered severe and permanent emotional injuries, including pain and suffering. Plaintiff also
10 has a substantial fear of suffering additional and ongoing harms, including but not limited to now
11 being at an increased risk of cancer, and future symptoms and harms associated with her autoimmune
12 disease and other injuries caused by Gardasil.

380. As a direct and proximate result of her Gardasil-induced injuries, Plaintiff has
suffered and continues to suffer economic losses, including considerable financial expenses for
medical care and treatment, and diminished income capacity, and she will continue to incur these
losses and expenses in the future.

381. Merck's conduct, as described above, was aggravated, oppressive, fraudulent, and 17 malicious. Merck regularly risks the lives of patients, including Plaintiff, with full knowledge of the 18 limited efficacy of Gardasil and the severe and sometimes fatal dangers of Gardasil. Merck has made 19 conscious decisions to not warn, or inform the unsuspecting public, including Plaintiff, and her 20 medical providers. Merck's conduct, including its false promotion of Gardasil and its failure to issue 21 appropriate warnings concerning the severe risks of Gardasil, created a substantial risk of significant 22 harm to children and patients who were being injected with Gardasil, and therefore warrants an award 23 of punitive damages. 24

382. WHEREFORE, Plaintiff requests that the Court enter judgment in her favor for
compensatory damages and punitive damages, together with interest, and costs herein incurred, and
all such other and further relief as this Court deems just and proper. Plaintiff also demands a jury trial
on the issues contained herein.

1	COUNT TWO							
2	STRICT LIABILITY							
3	(FAILURE TO WARN)							
4	383. Plaintiff incorporates by reference all other paragraphs of this Complaint as if fully set							
5	forth herein, and further alleges:							
6	384. Plaintiff brings this strict liability claim against Merck for failure to warn.							
7	385. At all times relevant to this litigation, Merck engaged in the business of researching,							
8	testing, developing, manufacturing, marketing, selling, distributing, and promoting Gardasil, which is							
9	defective and unreasonably dangerous to consumers, including Plaintiff, because it does not contain							
10	adequate warnings or instructions concerning the dangerous characteristics of Gardasil and its							
11	ingredients and adjuvants. These actions were under the ultimate control and supervision of Merck.							
12	386. Merck researched, developed, tested, manufactured, inspected, labeled, distributed,							
13	marketed, promoted, sold, and otherwise released into the stream of commerce Gardasil, and in the							
14	course of same, directly advertised or marketed the vaccine to consumers and end users, including							
15	Plaintiff and her medical providers, and Merck therefore had a duty to warn of the risks associated							
16	with the reasonably foreseeable uses of Gardasil and a duty to instruct on the proper,							
17	safe use of these products.							
18	387. At all times relevant to this litigation, Merck had a duty to properly research, test,							
19	manufacture, inspect, package, label, market, promote, sell, distribute, provide proper warnings, and							
20	take such steps as necessary to ensure that Gardasil did not cause users and consumers to suffer from							
21	unreasonable and dangerous risks. Merck had a continuing duty to instruct on the proper, safe use of							
22	these products. Merck, as manufacturer, seller, or distributor of vaccines, is held to the knowledge of							
23	an expert in the field.							
24	388. At the time of manufacture, Merck could have provided warnings or instructions							
25	regarding the full and complete risks of Gardasil because it knew or should have known of the							
26	unreasonable risks of harm associated with the use of and/or exposure to these products.							
27	389. At all times relevant to this litigation, Merck failed to properly investigate, study,							
28	research, test, manufacture, label or promote Gardasil. Merck also failed to minimize the dangers to							

children, patients, and consumers of Gardasil products and to those who would foreseeably use or be
 harmed by Gardasil, including Plaintiff.

390. Despite the fact that Merck knew or should have known that Gardasil posed a grave and 3 unreasonable risk of harm (including but not limited to increased risk of autoimmune disease, and the 4 various other Gardasil induced injuries that Plaintiff has sustained), it failed to warn of the risks 5 associated with Gardasil. The dangerous propensities of Gardasil and the carcinogenic characteristics 6 and autoimmune-inducing characteristics of Gardasil, as described in this Complaint, were known to 7 Merck, or scientifically knowable to Merck through appropriate research and testing by known 8 methods, at the time it distributed, supplied, or sold Gardasil, and not known to end users and 9 consumers, such as Plaintiff and her medical providers. 10

391. Merck knew or should have known that Gardasil and its ingredients and adjuvants
created significant risks of serious bodily harm to children and patients, as alleged herein, and Merck
failed to adequately warn patients, parents, medical providers and reasonably foreseeable users of the
risks and lack of efficacy of Gardasil. Merck has wrongfully concealed information concerning
Gardasil's dangerous nature and lack of efficacy and has further made false and misleading statements
concerning the safety and efficacy of Gardasil.

At all times relevant to this litigation, Merck's Gardasil products reached the intended
consumers, handlers, and users or other persons coming into contact with these products throughout
the United States, including Plaintiff, without substantial change in their condition as manufactured,
sold, distributed, labeled, and marketed by Merck.

21 393. Plaintiff was injected with Gardasil in its intended or reasonably foreseeable manner
 22 without knowledge of its unreasonable dangerous and inefficacious characteristics.

394. Plaintiff could not have reasonably discovered the defects and risks associated with
Gardasil before or at the time of her injection(s). Plaintiff relied upon the skill, superior knowledge,
and judgment of Merck.

395. Merck knew or should have known that the warnings disseminated with Gardasil were
 inadequate, and failed to communicate adequate information concerning the true risks and lack of
 efficacy of Gardasil and failed to communicate warnings and instructions that were appropriate and

adequate to render the products safe for their ordinary, intended, and reasonably foreseeable uses,
 including injections in teenagers.

396. The information that Merck did provide or communicate failed to contain relevant 3 warnings, hazards, and precautions that would have enabled patients, parents of patients and the 4 medical providers of patients to properly utilize, recommend or consent to the utilization of Gardasil. 5 Instead, Merck disseminated information that was inaccurate, false, and misleading and which failed 6 to communicate accurately or adequately the lack of efficacy, comparative severity, duration, and 7 extent of the serious risk of injuries associated Gardasil; continued to aggressively promote the 8 efficacy and safety of its products, even after it knew or should have known of Gardasil's 9 unreasonable risks and lack of efficacy; and concealed, downplayed, or otherwise suppressed, through 10 aggressive marketing and promotion, any information or research about the risks, defects and dangers 11 of Gardasil. 12

397. To this day, Merck has failed to adequately and accurately warn of the true risks of
Plaintiff's injuries, including but not limited to, autoimmune diseases, including POTS and
dysautonomia, associated with the use of and exposure to Gardasil, and has failed to warn of the
additional risks that Plaintiff is now exposed to, including, but not limited to, the increased risk of
cancer, and other potential side effects and ailments.

398. As a result of Merck's failure to warn and false promotion, Gardasil is and was
defective and unreasonably dangerous when it left the possession and/or control of Merck, was
distributed by Merck, and used by Plaintiff.

399. Merck is liable to Plaintiff for injuries caused by its failure, as described above, to
 provide adequate warnings or other clinically relevant information and data regarding Gardasil, the
 lack of efficacy and serious risks associated with Gardasil and its ingredients and adjuvants.

400. The defects in Merck's Gardasil vaccine were substantial and contributing factors in
causing Plaintiff's injuries, and, but for Merck's misconduct and omissions and Gardasil's defects,
including its defective labeling and false promotion, Plaintiff would not have sustained her injuries
which she has sustained to date, and would not have been exposed to the additional prospective risk
and dangers that are associated with Gardasil.

Had Merck not engaged in the negligent and fraudulent conduct alleged herein and/or 1 401. 2 had Merck, via its labeling, advertisements, and promotions provided adequate and truthful warnings and properly disclosed and disseminated the true risks, limitations, and lack of efficacy associated 3 with Gardasil to medical providers, patients and the public, then upon information and belief, 4 Plaintiff's medical providers would not have offered or recommended Gardasil to Plaintiff. 5 Moreover, even if after Merck's dissemination of truthful information concerning the true risks and 6 efficacy limitation of Gardasil, Plaintiff's medical providers had offered Gardasil, then upon 7 information and belief, the providers would have heeded any warnings issued by Merck and relayed to 8 Plaintiff the safety risks and efficacy limitations that Merck should have warned her about, but failed 9 to do so. Had Plaintiff been informed of the true risks and efficacy limitation concerning Gardasil, 10 through her medical providers or through Merck's ubiquitous direct-to-consumer promotional 11 marketing, on which she relied, then Plaintiff would not have consented to being injected with 12 Gardasil. 13

402. As a proximate result of Merck's wrongful acts and omissions and its negligent and fraudulent testing, labeling, manufacturing, and promotion of Gardasil, Plaintiff has suffered and continues to suffer severe and permanent physical injuries, including, but not limited to, her autoimmune disease and associated symptomology and has suffered severe and permanent emotional injuries, including pain and suffering. Plaintiff also has a substantial fear of suffering additional and ongoing harms, including but not limited to now being at an increased risk of cancer, and future symptoms and harms associated with her autoimmune disease and other injuries caused by Gardasil.

403. As a direct and proximate result of her Gardasil-induced injuries, Plaintiff has suffered
and continues to suffer economic losses, including considerable financial expenses for medical care
and treatment, and diminished income capacity and she will continue to incur these losses and
expenses in the future.

404. Merck's conduct, as described above, was oppressive, fraudulent, and malicious.
Merck regularly risks the lives of teenagers, including Plaintiff, with full knowledge of the limited
efficacy of Gardasil and the severe and sometimes fatal dangers of Gardasil. Merck has made
conscious decisions to not warn or inform the unsuspecting public, including Plaintiff and her medical

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providers. Merck's conduct, including its false promotion of Gardasil and its failure to issue
 appropriate warnings concerning the severe risks of Gardasil, created a substantial risk of significant
 harm to children, teenagers, and patients who were being injected with Gardasil, and therefore
 warrants an award of punitive damages.

5 405. WHEREFORE, Plaintiff requests that the Court enter judgment in her favor for all 6 compensatory and punitive damages, together with interest, and costs herein incurred, and all such 7 other and further relief as this Court deems just and proper. Plaintiff also demands a jury trial on the 8 issues contained herein.

COUNT THREE

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STRICT LIABILITY

(MANUFACTURING DEFECT)

406. Plaintiff incorporates by reference all other paragraphs of this Complaint as if fully set
forth herein, and further alleges:

407. Plaintiff brings this strict liability claim against Merck for manufacturing defect.
408. At all times relevant to this litigation, Merck engaged in the business of researching,
testing, developing, manufacturing, marketing, selling, distributing, and promoting Gardasil, which is
defective and unreasonably dangerous to consumers, including Plaintiff, because of manufacturing
defects, which patients, including Plaintiff and her medical providers did not expect.

409. Upon information and belief, the Gardasil vaccines injected into Plaintiff were defective
and unreasonably dangerous because they failed to comply with manufacturing specifications required
by the governing manufacturing protocols and also required by the regulatory agencies, including but
not limited to the FDA, by among other things, containing ingredients and toxins that were not
disclosed in the FDA-approved specifications and/or otherwise not disclosed in the package insert.

410. Upon information and belief, and as way of example, the Gardasil injected into Plaintiff
was defective and unreasonably dangerous because it failed to comply with the approved
manufacturing specifications, by containing dangerous and undisclosed HPV L1-DNA fragments, and
these DNA fragments could act as a Toll-Like Receptor 9 (TLR9) agonist, further adjuvanting the
vaccine and making it more potent and dangerous than intended.

411. Upon information and belief, and as way of example, the Gardasil injected into Plaintiff
 was defective and unreasonably dangerous because it failed to comply with the approved
 manufacturing specifications, by containing dangerous and undisclosed ingredients and neurotoxins,
 including but not limited to, phenylmethylsulfonyl fluoride (PMSF), a toxic nerve agent that is not
 intended for human consumption or injections.

412. At all times relevant to this litigation, Merck's Gardasil products reached the intended
consumers, handlers, and users or other persons coming into contact with these products throughout
the United States, including Plaintiff, without substantial change in their condition as designed,
manufactured, sold, distributed, labeled, and marketed by Merck.

413. Plaintiff and her medical providers could not reasonably have discovered the defects,
including the manufacturing defects, and risks associated with Gardasil before or at the time of her
injection(s). Plaintiff relied upon the skill, superior knowledge, and judgment of Merck.

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414. Merck is liable to Plaintiff for injuries caused as a result of its manufacturing defects.

4 415. The defects in Merck's Gardasil vaccine were substantial and contributing factors in
causing Plaintiff's injuries, and, but for Merck's misconduct and omissions and Gardasil's defects,
including but not limited to its manufacturing defects, Plaintiff would not have sustained the injuries
she has sustained to date, and would not have been exposed to the additional prospective risk and
dangers associated with Gardasil.

416. As a proximate result of Merck's wrongful acts and Gardasil's manufacturing defects,
Plaintiff has suffered and continues to suffer severe and permanent physical injuries and associated
symptomology and has suffered severe and permanent emotional injuries, including pain and
suffering. Plaintiff also has a substantial fear of suffering additional and ongoing harms, including but
not limited to now being at an increased risk of cancer, and future symptoms and harms associated
with her autoimmune disease and other injuries caused by Gardasil.

417. As a direct and proximate result of her Gardasil-induced injuries, Plaintiff has
suffered and continues to suffer economic losses, including considerable financial expenses for
medical care and treatment, and diminished income capacity, and she will continue to incur these
losses and expenses in the future.

1 418. Merck's conduct, as described above, was oppressive, fraudulent, and malicious. 2 Merck regularly risks the lives of patients, including Plaintiff, with full knowledge of the limited 3 efficacy of Gardasil and the severe and sometimes fatal dangers of Gardasil. Merck has made conscious decisions to not warn, or inform the unsuspecting public, including Plaintiff, and her 4 medical providers. Merck's conduct, including its false promotion of Gardasil and its failure to issue 5 appropriate warnings concerning the severe risks of Gardasil, created a substantial risk of significant 6 7 harm to children and patients who were being injected with Gardasil, and therefore warrants an award 8 of punitive damages.

9 419. WHEREFORE, Plaintiff requests that the Court enter judgment in her favor for
10 compensatory and punitive damages, together with interest, and costs herein incurred, and all such
11 other and further relief as this Court deems just and proper. Plaintiff also demands a jury trial on the
12 issues contained herein.

COUNT FOUR

BREACH OF EXPRESS WARRANTY

420. Plaintiff incorporates by reference all other paragraphs of this Complaint as if fully set
forth herein, and further alleges:

421. Merck engaged in the business of testing, researching, manufacturing, labeling,
marketing, selling, distributing, and promoting Gardasil, which is defective and unreasonably
dangerous to consumers, including Plaintiff.

422. At all times relevant to this litigation, Merck expressly represented and warranted 20 through statements made in its Gardasil label, publications, television advertisements, billboards, print 21 advertisements, online advertisements and website, and other written materials intended for 22 consumers, patients, parents of minor-aged patients, medical providers and the general public, that 23 Gardasil was safe and effective at preventing cancer. Merck advertised, labeled, marketed, and 24 promoted Gardasil, representing the quality to consumers, patients, medical providers and the public 25 in such a way as to induce their purchase or use, thereby making an express warranty that Gardasil 26 would conform to the representations. 27

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423. These express representations included incomplete warnings and instructions that
purport, but fail, to include the complete array of risks associated with Gardasil. Merck knew and/or
should have known that the risks expressly included in Gardasil's promotional material and labels did
not and do not accurately or adequately set forth the risks of developing the serious injuries
complained of herein. Nevertheless, Merck falsely and expressly represented that Gardasil was "safe"
for use by individuals such as Plaintiff, and/or that Gardasil was "effective" in preventing cancer and
that anyone who was vaccinated with Gardasil would be "one less" person with cancer.

8 424. The representations about Gardasil, as set forth herein, contained or constituted
9 affirmations of fact or promises made by the seller to the buyer, which related to the goods and
10 became part of the basis of the bargain, creating an express warranty that the goods would conform to
11 the representations.

425. Merck breached these warranties because, among other things, Gardasil is ineffective at
preventing cancer, defective, dangerous, unfit for use, and is associated with a myriad of dangerous
and undisclosed risks, including, but not limited to, the risk of autoimmune disease, including POTS,
the risk of developing cervical cancer in women (even though Merck promoted it as preventing
cervical cancer), and the risk of fertility problems for young girls. Specifically, Merck breached the
warranties in the following ways:

Representing to patients and the medical community, including Plaintiff, her a) 18 parents and/or her medical providers that Gardasil is effective in preventing 19 cancer, including anal and cervical cancer, when Merck knew that contrary to 20 these representations (i) no clinical studies were performed to test if Gardasil 21 prevents cancer; (ii) the clinical studies confirmed that Gardasil is indeed 22 ineffective when used in patients who have previously been exposed to HPV, 23 and that Gardasil actually increases the risk of cancer in a patient who has been 24 previously exposed to HPV; and (iii) there are safer and more effective methods 25 of monitoring for and attempting to prevent cervical or anal cancer, including 26 but not limited to regular testing, such as regular Pap smears for cervical cancer, 27 and monitoring for anal cancer. 28

 b) Representing to patients and the medical community, including Plaintiff and her medical providers that Gardasil is safe, when in reality, Gardasil causes and presents serious risks of cancer, autoimmune disease, including but not limited to POTS, and other grave illnesses as outlined herein;

c) Engaging in false advertising and disease mongering by scaring parents and teenagers into believing that cervical and anal cancer is far more prevalent than it really is; that all cervical and anal cancer was linked to HPV; that Gardasil prevented cervical cancer, when in reality none of these representations were true as cervical cancer rates were declining in the United States due to Pap testing and Gardasil has not been shown to prevent against all strains of HPV that are associated with cervical cancer and indeed it has never been shown to prevent cervical or anal cancer.

426. Merck had sole access to material facts concerning the nature of the risks and defects
associated with Gardasil as expressly stated within its promotional material and labels, and Merck
knew that patients and users such as Plaintiff could not have reasonably discovered the truth about the
inefficacies and serious risks associated with Gardasil as alleged herein.

17 427. Plaintiff had no knowledge of the falsity or incompleteness of Merck's statements and
18 representations concerning Gardasil.

428. Plaintiff was exposed to and relied upon the ubiquitous promotional material and 19 representations Merck made in its direct-to-consumer advertisements and marketing materials 20 concerning the safety and efficacy of Gardasil, including: that Gardasil prevents cervical and anal 21 cancer and these cancers are prevalent (even though children rarely get cervical or anal cancer and Pap 22 tests are the best frontline defense in detecting and fighting cervical cancer); that "good mothers" 23 vaccinate their children and that Gardasil is perfectly safe. However, had Merck in these 24 advertisements not engaged in disease mongering and deception, but instead had informed her the 25 truth about the serious risks of Gardasil (as outlined in this Complaint) and its lack of efficacy, she 26 would never have consented to being injected with Gardasil, nor would Plaintiff have consented to the 27

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Gardasil injection(s) had she been adequately informed about the questionable efficacy and serious
 risks associated with Gardasil.

429. As a proximate result of Merck's wrongful acts and breaches of warranties concerning
the safety and efficacy of Gardasil, Plaintiff has suffered and continues to suffer severe and permanent
physical injuries, and associated symptomology and has suffered severe and permanent emotional
injuries, including pain and suffering. Plaintiff also has a substantial fear of suffering additional and
ongoing harms, including but not limited to now being at an increased risk of cancer, and future
symptoms and harms associated with her autoimmune disease and other injuries caused by Gardasil.

9 430. As a direct and proximate result of her Gardasil-induced injuries, Plaintiff has
10 suffered and continues to suffer economic losses, including considerable financial expenses for
11 medical care and treatment, and diminished income capacity and she will continue to incur these
12 losses and expenses in the future.

431. Merck's conduct, as described above, was oppressive, fraudulent, and malicious. 13 Merck regularly risks the lives of patients, including Plaintiff, with full knowledge of the limited 14 efficacy of Gardasil and the severe and sometimes fatal dangers of Gardasil. Merck has made 15 conscious decisions to not warn, or inform the unsuspecting public, including Plaintiff and her 16 medical providers. Merck's conduct, including its false promotion of Gardasil and its failure to issue 17 appropriate warnings concerning the severe risks of Gardasil, created a substantial risk of significant 18 harm to children and patients who were being injected with Gardasil, and therefore warrants an award 19 of punitive damages. 20

432. WHEREFORE, Plaintiff requests that the Court enter judgment in her favor for
compensatory and punitive damages, together with interest, and costs herein incurred, and all such
other and further relief as this Court deems just and proper. Plaintiff also demands a jury trial on the
issues contained herein.

COUNT FIVE

COMMON LAW FRAUD

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433. Plaintiff incorporates by reference all other paragraphs of this Complaint as if fully set
forth herein, and further alleges:

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434. Merck is the researcher, manufacturer, labeler, and promoter of Gardasil.

435. Merck marketed Gardasil to and for the benefit of patients, including teenagers such as
Plaintiff and her medical providers.

4 436. Merck had a duty to deal honestly and truthfully with regulators, patients, consumers
5 and medical providers in its development, testing, marketing, promotion, and sale of Gardasil.

437. Merck's duty of care owed to patients and medical providers included providing
accurate, complete, true, and correct information concerning the efficacy and risks of Gardasil in its
direct-to-consumer advertisements, promotional material, and labeling.

9 438. At all times relevant to this litigation, Merck knew or should have known of the hazards
10 and dangers of Gardasil and specifically, the serious, debilitating and potentially fatal adverse events
11 associated with Gardasil, including but not limited to autoimmune diseases, increased risk of cancer,
12 and death.

439. At all times relevant to this litigation, Merck knew or should have known that its poorly
designed clinical trials and studies were insufficient to test the true long-term safety and efficacy of
Gardasil.

440. At all times relevant to this litigation, Merck expressly represented through statements it
made in its publications, ubiquitous television advertisements, billboards, print advertisements, online
advertisements and website, and other written materials intended for consumers, patients, parents of
minor-aged patients, medical providers and the general public, that Gardasil was safe and effective at
preventing cancer.

441. These express representations included incomplete warnings and instructions that
purport, but fail, to include the complete array of risks associated with Gardasil. By way of example
Merck's marketing material, including its "One Less" television and print advertisement campaign
(including but not limited to Gardasil posters in medical facilities and doctors' offices), which
Plaintiff had been exposed to, stated that Gardasil was safe, that Gardasil was effective in preventing
cancer, that Gardasil was a "cervical cancer vaccine," and that any young child or teenager who was
vaccinated with Gardasil would lead to "one less" person with cervical or anal cancer. The only safety

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warnings Merck provided in these marketing materials was that a patient could get pain, swelling or
 redness at injection site, fever, and/or nausea.

3 442. The ubiquitous nature of these Gardasil commercials and the Gardasil marketing
4 campaign gave the impression that cervical cancer was on the rise and more prevalent than it actually
5 was, and that all good mothers vaccinate their children with the "cervical cancer vaccine."

6 443. Merck knew or should have known that the risks expressly included in Gardasil's
7 promotional material and labels did not and do not accurately or adequately set forth the true and
8 complete risks of developing the serious injuries that are associated with Gardasil, as previously
9 alleged herein, and which include but are not limited to POTS, systemic adverse events, autoimmune
10 disease, increased risk of cancer, and death.

444. The same promises of efficacy and limited and incomplete warnings Merck relayed in
its direct-to-consumer advertising, were what Plaintiff's medical providers relayed to her when they
recommended Gardasil – i.e., that if Plaintiff got vaccinated with Gardasil, it would prevent cancer.
Plaintiff was not warned of any potential adverse reactions to the Gardasil vaccine by her healthcare
provider.

445. Plaintiff had been exposed to Merck's marketing material concerning Gardasil, 16 including the aforementioned "One Less" marketing campaign and other print advertisements and 17 posters at doctors' offices, and the representations made by Merck therein that Gardasil is effective at 18 preventing cervical and anal cancer, that Gardasil is safe and that its only side-effects are essentially 19 minor injection site pain and swelling, and the possible onset of a fever or nausea. Prior to providing 20 consent to inject Plaintiff with the Gardasil vaccine, Plaintiff was never informed by Merck, or 21 anyone else, that Gardasil is linked to a host of serious debilitating and chronic adverse events 22 including, autoimmune diseases (including, but not limited to, POTS), increased risk of cancer, and 23 death. 24

446. Prior to providing consent to inject Plaintiff with the Gardasil vaccine, Plaintiff was
 never informed by Merck, or anyone else, that Merck had not conducted the proper testing necessary
 to demonstrate the efficacy and full safety of Gardasil.

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447. Prior to providing consent to inject Plaintiff with the Gardasil vaccine, Plaintiff was
 never informed by Merck, or anyone else, that Merck had, as alleged herein, manipulated its clinical
 studies to mask and conceal the adverse events associated with Gardasil.

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448. Prior to providing consent to inject Plaintiff with the Gardasil vaccine, Plaintiff was never informed by Merck, or anyone else, that the Gardasil clinical trials never established that Gardasil can prevent cervical or anal cancer, even though Merck in its promotional material falsely represented that Gardasil was a "cervical cancer vaccine" and that a patient who received Gardasil would result in "one less" woman or man getting cancer.

9 449. Merck's representations were false, because in truth, Gardasil has not been proven to
10 prevent cervical or anal cancer and is associated with a myriad of dangerous and undisclosed risks,
11 including, but not limited to, the risk of autoimmune disease, including POTS, increased risk of
12 developing cancer, and other serious side effects. The false representations Merck made to the
13 patients, children, teenagers, the parents of children and teenagers, the medical community, including
14 to Plaintiff, included:

that Gardasil is effective in preventing cervical and anal cancer, when Merck a) 15 knew that, contrary to these representations (i) no clinical studies were 16 performed to test whether Gardasil prevents cancer; and (ii) the clinical studies 17 confirmed that Gardasil is indeed ineffective when used in patients who have 18 previously been exposed to HPV, and that Gardasil actually increases the risk of 19 cervical cancer in any child or patient who has been previously exposed to HPV; 20 that Gardasil is safe, when in reality, Gardasil causes and presents severe risks **b**) 21 of cancer (including cervical cancer, the very cancer it is promoted as 22 preventing), fertility problems, autoimmune disease, including POTS, OI, and 23 other grave illnesses; 24 false advertising and disease mongering by scaring parents into believing that c) 25 cervical and anal cancer were far more prevalent than it really was; that Gardasil 26 prevented cervical and anal cancer; and that Gardasil only had risks of injection 27

> 75 COMPLAINT

site pain and fever, when in reality none of these representations were true as

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cervical cancer rates were declining in the United States due to Pap testing and Gardasil has not been shown to prevent cervical or anal cancer, and indeed some studies demonstrated that it actually increased the risk of cervical cancer; and Gardasil was linked to a host of serious, chronic and sometimes fatal diseases, including autoimmune diseases, as previously outlined in this Complaint.

6 450. These representations and other similar representations were made by Merck to the
7 public, including to Plaintiff, with the intent that parents would either seek out Gardasil from their
8 medical providers or otherwise would provide their consent when they were offered Gardasil.

9 451. At the time she provided her consent to the Gardasil injection(s), Plaintiff's mother was
10 not aware of the falsity of Merck's aforementioned representations concerning the safety and efficacy
11 of Gardasil.

452. Plaintiff reasonably and justifiably relied upon the truth of the assurance made by
Merck in its direct-to-consumer marketing concerning the efficacy and safety of Gardasil (which were
also echoed by Plaintiff's medical providers), when her mother provided consent for her to be injected
with the Gardasil vaccine.

453. Had Merck's advertisements and promotional material, which Merck targeted to
teenagers and the parents of teenagers, and which Plaintiff received and on which she relied, provided
complete and truthful warnings and properly disclosed and disseminated the true risks, limitations and
lack of efficacy associated with Gardasil, then Plaintiff would not have consented to being injected
with Gardasil.

454. Merck also engaged in a number of additional fraudulent activities that led to regulators,
medical providers (upon information and belief, including but not limited Plaintiff's medical
providers), and the general public (including directly and/or indirectly Plaintiff) to be duped into
believing that Gardasil is safe and effective. These fraudulent acts are outlined in greater detail in the
preceding paragraphs of this Complaint, and included, among others:

 a) Failing to test Gardasil against a true inert placebo and lying to the public that Gardasil was tested against a placebo, when in reality, all, or nearly all, studies used a toxic placebo that included the dangerous aluminum adjuvant AAHS.

b)	Failing to conduct a sufficient number of studies for the targeted patient
	population which included pre-teen girls (and boys) between the ages of nine
	and 12.

- c) Not using the commercial dosage (and instead using a lower dosage of the adjuvant and ingredients) in one of the key clinical trials, which was used to obtain licensing for the commercial dosage of Gardasil;
- d) Using very restrictive exclusionary criteria in the clinical study patient population (including for example, exclusion of anyone who had prior abnormal Pap tests, who had a history of immunological or nervous system disorders or was allergic to aluminum or other ingredients), but then not revealing or warning about these exclusionary criteria in the label and knowing that for most of these ingredients and allergies, there are limited resources for the public to test for such allergies in advance of being vaccinated;
 - e) Failing to disclose all of the ingredients in Gardasil, including but not limited to the fact that Gardasil contains dangerous HPV L1-DNA fragments and that these DNA fragments could act as a Toll-Like Receptor 9 (TLR9) agonist – further adjuvanting the vaccine and making it more potent and dangerous.

455. Merck engaged in the above mentioned fraudulent conduct as well as the additional
fraudulent conduct detailed throughout this Complaint with the intent to enhance Gardasil's safety and
efficacy profile and to conceal Gardasil's serious risks and efficacy shortcomings in order to secure
regulatory approval and more importantly, so as to encourage physicians and medical providers to
recommend Gardasil to patients and to prepare and encourage patients to request and consent to
Gardasil injections.

456. Plaintiff could not reasonably have discovered the falsity of Merck's representations,
the fraudulent nature of Merck's conduct, and the defects and risks associated with Gardasil before or
at the time of her injection(s). Plaintiff relied upon the skill, superior knowledge, and judgment of
Merck, the manufacturer, labeler, and promoter of Gardasil, and they detrimentally relied upon
Merck's fraudulent, false, and misleading statements, omissions, and conduct.

457. As a proximate result of Merck's fraudulent, false, and misleading statements,
omissions, and conduct concerning the safety and efficacy of Gardasil, Plaintiff has suffered and
continues to suffer severe and permanent physical injuries, and associated symptomology and has
suffered severe and permanent emotional injuries, including pain and suffering. Plaintiff also has a
substantial fear of suffering additional and ongoing harms, including but not limited to now being at
an increased risk of cancer, and future symptoms and harms associated with her autoimmune disease
and other injuries caused by Gardasil.

8 458. As a direct and proximate result of her Gardasil-induced injuries, Plaintiff has
9 suffered and continues to suffer economic losses, including considerable financial expenses for
10 medical care and treatment, and diminished income capacity and she will continue to incur these
11 losses and expenses in the future.

12 459. Merck's conduct, as described above, was oppressive, fraudulent, and malicious.
13 Merck regularly risks the lives of patients, including Plaintiff, with full knowledge of the limited
14 efficacy of Gardasil and the severe and sometimes fatal dangers of Gardasil. Merck has made
15 conscious decisions to not warn, or inform the unsuspecting public, including Plaintiff and her
16 medical providers. Merck's conduct, including its false promotion of Gardasil and its failure to issue
17 appropriate warnings concerning the severe risks of Gardasil, created a substantial risk of significant
18 harm to children and patients who were being injected with Gardasil.

460. WHEREFORE, Plaintiff requests that the Court enter judgment in her favor for
compensatory and punitive damages, together with interest, and costs herein incurred, and all such
other and further relief as this Court deems just and proper. Plaintiff also demands a jury trial on the
issues contained herein.

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VIOLATION OF WEST VIRGINIA'S CONSUMER CREDIT AND PROTECTION ACT

COUNT SIX

465. Plaintiff incorporates by reference all other paragraphs of this Complaint as if fully set
forth herein, and further alleges:

466. West Virginia's Consumer Credit and Protection Act ("WVCCPA"), W. Va. Code §
46A-6-101, et seq., protects both consumers and competitors by promoting fair competition in

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commercial markets for goods and services. West Virginia's Consumer Credit and Protection Act is
 interpreted broadly and provides a cause of action for any unlawful, unfair, or fraudulent business act
 or practice. Any unlawful, unfair, or fraudulent business practice that causes injury to consumers falls
 within the ambit of West Virginia's Consumer Credit and Protection Act.

467. The WVCCPA contains an all-encompassing, blanket prohibition against "[u]nfair
methods of competition and unfair or deceptive acts or practices in the conduct of any trade or
commerce...." W. VA. Code § 46A-6-104. The WVCCPA delineates at least 15 types of conduct that
constitute per se violations. W. Va. Code § 46A-6-102(7). The statutory list is not intended to be all
inclusive. *Id.*

468. The WVCCPA specifically prohibits "The act, use or employment by any person of any
deception, fraud, false pretense, false promise or misrepresentation, or the concealment, suppression
or omission of any material fact with intent that others rely upon such concealment, suppression or
omission, in connection with the sale or advertisement of any goods or services, whether or not any
person has in fact been misled, deceived or damaged thereby...." as a delineated, per se violation. W.
Va. Code § 46A-6-102(7)(M).

469. Merck engaged in substantial advertising and marketing of Gardasil within the State of
West Virginia.

18 470. Because of Merck's unlawful, fraudulent, and unfair business practices, Plaintiff was
19 misled into purchasing and consenting to the Gardasil injection.

471. As set forth in the preceding paragraphs, Defendants have engaged in the unlawful
business practice of misleading Plaintiff regarding the Gardasil vaccines' true safety. Defendants'
deceptive and unlawful marketing practices have violated numerous West Virginia laws, including,
inter alia: W. Va. Code § 46A-6-107 (breach of express warranty); W. Va. Code §§ 46A-6-102; 46A6-104 (false advertising and marketing); and W. Va. Code § 46A-6-101, et seq. (violations of West
Virginia's Consumer Credit and Protection Act).

26 472. Merck widely advertised and promoted Gardasil as a safe and effective vaccine that had
 27 no serious side effects.

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473. Yet, contrary to its above referenced false claims concerning the safety and efficacy of

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Gardasil, Merck knew, or should have known, that Gardasil was ineffective, unreasonably dangerous
 and defective, and had a propensity to cause serious and life-threatening side effects, including but not
 limited to autoimmune diseases and other grave injuries as outlined in this Complaint.

4 474. The false, deceptive, and misleading actions, statements, and representations made by
5 Merck, as alleged in this Complaint, are unlawful, fraudulent, and unfair business practices and acts
6 within the meaning of the WVCCPA. See e.g., W. Va. Code § 46A-6-102, et seq.

475. Merck's concealment of the autoimmune risks and other adverse events outlined in this
Complaint was a material omission that consumers, patients, parents, and prescribing healthcare
professionals should have known about prior to purchasing, consenting to injections of, or prescribing
Gardasil.

476. Merck's concealment of the lack of efficacy and false representations concerning the
efficacy of Gardasil in preventing cancer was a material false representation and omission that
consumers, patients, parents, and prescribing healthcare professionals should have known about prior
to purchasing, consenting to injections of, or prescribing Gardasil.

477. Merck had sole access to material facts concerning the nature of the risks and defects
associated with Gardasil as expressly stated within its promotional material and labels, and Merck
knew that patients and users such as Plaintiff and her medical providers could not have reasonably
discovered the truth about the inefficacies and serious risks associated with Gardasil as alleged herein.

19 478. Plaintiff had no knowledge of the falsity or incompleteness of Merck's statements and
 20 representations concerning Gardasil.

479. Plaintiff reasonably and justifiably relied upon the truth of the assurance made by
Merck in its direct-to-consumer marketing concerning the efficacy and safety of Gardasil (which were
also echoed by Plaintiff's medical providers), when her mother provided her consent to being injected
with the Gardasil vaccine.

480. Had Merck's advertisements and promotional material, which Merck targeted to
teenagers and the parents of teenagers, and which Plaintiff received and on which she relied, provided
complete and truthful warnings and properly disclosed and disseminated the true risks, limitations,

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and lack of efficacy associated with Gardasil, then Plaintiff would never have consented to being
 injected with Gardasil.

481. As a direct and proximate result of Merck's unlawful, fraudulent, and unfair business
practices, Plaintiff has sustained injuries and economic damages as outlined herein, including but not
limited to, agreeing to being injected with Gardasil, which upon information and belief, costs more
than \$100 per vile.

482. As a result of Merck's violation of the WVCCPA, Plaintiff seeks an order of this Court
enjoining Merck from continuing these unlawful, fraudulent, and unfair practices and awarding
Plaintiff remedies, including but not limited to disgorgement of Merck's profits, restitution, fees, and
all other remedies available under law.

11 483. WHEREFORE, Plaintiff requests that the Court enter judgment in her favor for 12 restitution, disgorgement of Merck's ill-gotten profits, punitive damages, and all other permissible 13 monetary relief, together with interest, costs herein incurred, attorney fees, and all such other and 14 further relief as this Court deems just and proper. Plaintiff also requests that the Court issue an 15 injunction prohibiting Merck from continuing its false advertising and unlawful acts and practices 16 concerning Gardasil and to grant any other preliminary or permanent equitable relief as deemed 17 appropriate.

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PRAYER FOR RELIEF

WHEREFORE, Plaintiff, ELIZABETH LANDERS on behalf of her minor child I.L, requests
that the Court enter judgment in her favor and against Merck & Co., Inc., and Merck, Sharp and
Dohme Corporation (collectively "Merck") as to all causes of action, and awarding as follows:
A. For compensatory damages, in an amount exceeding this Court's jurisdictional

- minimum and to be proven at trial;
- B. For economic and non-economic damages in an amount to be proven at trial;
- C. For medical, incidental, hospital, psychological and other expenses in an amount to be proven at trial;
- D. For loss of earnings and earnings capacity, in an amount to be proven at trial;
- E. For an award of pre-judgment and post-judgment interest as provided by law;

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1	F. For exemplary and punitive damages against Merck;						
2	G. For preliminary and/or permanent injunctive relief against Merck;						
3	H. For an award providing for payment of reasonable fees, court costs, and other litigation						
4	expenses as permitted by law;						
5	I. For such other and further relief as this Honorable Court may deem just and proper.						
6			DEM	AND FOR JURY	TRIAL		
7	Pursuant to	Rule 3	8(b) of the Fede	eral Rules of Civil	Procedure, Plaintiff, ELIZABETH		
8	LANDERS on behalf of her minor child I.L, hereby demands a jury trial on <i>all</i> of her claims, causes						
9	of action and issue	es that a	are triable by ju	·y.			
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11	Dated: April 1, 20)22		MORGAN &	MORGAN, P.A.		
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				By: <u>/s/ Mark</u>	-		
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24				Application for	r <i>pro hac vice</i> admission to be filed		
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