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IN THE UNITED STATES DISTRICT COURT  
FOR THE NORTHERN DISTRICT OF CALIFORNIA  
SAN FRANCISCO DIVISION

DOLORES GOULD,

Plaintiff,

vs.

JOHNSON & JOHNSON, and JOHNSON &  
JOHNSON CONSUMER COMPANIES, INC.,

Defendants.

Case No.

**COMPLAINT**

**JURY TRIAL DEMANDED**

**I. COMPLAINT**

Plaintiff Dolores Gould, by and through undersigned counsel, brings this action against Defendants Johnson & Johnson (“J&J) and Johnson & Johnson Consumer Companies, Inc. (“J&J Consumer”) as follows:

**II. INTRODUCTION**

1. This action arises out of Plaintiff Dolores Gould’s diagnosis of uterine cancer, which was directly and proximately caused by her regular and prolonged exposure to talcum powder, contained in Defendants’ Johnson & Johnson Baby Powder (hereinafter “J&J Baby Powder”) and Shower to Shower. Plaintiff brings this cause of action against Defendants for claims arising from the direct and proximate result of Defendants’ and/or their corporate predecessors’ negligent,

1 willful, and wrongful conduct in connection with the design, development, manufacture, testing,  
2 packaging, promoting, marketing, distribution, labeling, and/or sale of the products known as J&J  
3 Baby Powder and Shower to Shower (hereinafter collectively referred to as “Products”).

4 **III. PARTIES**

5 2. Plaintiff was born in 1975, and used J&J Baby Powder and Shower to Shower, the  
6 “Products,” for nearly her entire life. As a direct and proximate result of using the Products,  
7 Plaintiff was diagnosed with uterine cancer in 2006. Plaintiff resides in Oakley, in Contra Costa  
8 County, California. Plaintiff resided at the Great Lakes Naval Station, Great Lakes, Illinois at the  
9 time of her diagnosis.

10 3. Defendant, Johnson & Johnson (“J&J”), is a New Jersey corporation with its  
11 principal place of business in the State of New Jersey.

12 4. At all pertinent times, Johnson & Johnson was engaged in the business of  
13 manufacturing, marketing, testing, promoting, selling, and/or distributing the Products. At all  
14 pertinent times, Johnson & Johnson regularly transacted, solicited, and conducted business in all  
15 States of the United States, including the State of California.

16 5. Defendant, Johnson & Johnson Consumer Companies, Inc. is a New Jersey  
17 corporation with its principal place of business in the State of New Jersey.

18 6. At all pertinent times, Johnson & Johnson Consumer Companies, Inc. was engaged  
19 in the business of manufacturing, marketing, testing, promoting, selling, and/or distributing the  
20 Products. At all pertinent times, Johnson & Johnson regularly transacted, solicited, and conducted  
21 business in all States of the United States, including the State of California.

22 7. At all pertinent times, all Defendants were engaged in the research, development,  
23 manufacture, design, testing, sale and marketing of the Products, and introduced such products into  
24 interstate commerce with knowledge and intent that such products be sold in the State of California.

25 **IV. JURISDICTION AND VENUE**

26 8. This Court has original jurisdiction pursuant to 28 U.S.C. § 1332(d) because  
27 complete diversity exists between Plaintiff and Defendants, and the matter in controversy, exclusive  
28 of interest and costs, exceeds the sum or value of \$75,000.

1 9. This Court has personal jurisdiction over Defendants because Defendants are  
2 authorized to conduct and do conduct business in the State of California. Defendants have  
3 marketed, promoted, distributed, and sold the Products in the state of California and Defendants  
4 have sufficient minimum contacts with this State and/or sufficiently avail themselves of the markets  
5 in this State through their promotion, sales, distribution and marketing within this State to render  
6 the exercise of jurisdiction by this Court permissible.

7 10. Venue is proper in this Court pursuant to 28 U.S.C. §§ 1391(a) and (b) because a  
8 substantial part of the events or omissions giving rise to Plaintiff’s claims occurred in this judicial  
9 district. Venue is also proper under 18 U.S.C. § 1965(a) because Defendants transact substantial  
10 business in this District.

11 **V. FACTS COMMON TO ALL COUNTS**

12 **A. Background: Talc as a Carcinogen and Defendants’ Knowledge**

13 11. Talc is a magnesium trisilicate and is mined from the earth. Talc is an inorganic  
14 mineral.

15 12. Talc is the main substance in talcum powders. The Johnson & Johnson Defendants  
16 manufactured the Products. The Products are composed almost entirely of talc.

17 13. At all pertinent times, a feasible alternative to the Products has existed. For example,  
18 cornstarch is an organic carbohydrate that is quickly broken down by the body with no known  
19 health effects. Cornstarch powders have been sold and marketed for the same uses as the Products  
20 with nearly the same effectiveness.

21 14. Historically, “Johnson’s Baby Powder” has been a symbol of freshness, cleanliness,  
22 and purity. During the time in question, the Johnson & Johnson Defendants advertised and  
23 marketed this product as the beacon of “freshness” and “comfort”, eliminating friction on the skin,  
24 absorbing “excess wetness” helping keep skin feeling dry and comfortable, and “clinically proven  
25 gentle and mild.” The Johnson & Johnson Defendants instructed women through advertisements to  
26 dust themselves with this product to mask odors. The bottle of “Johnson’s Baby Powder”  
27 specifically targets women by stating, “For you, use every day to help feel soft, fresh, and  
28 comfortable.”

1           15. During the time in question, the Johnson & Johnson Defendants advertised and  
2 marketed the product “Shower to Shower” as safe for use by women as evidenced in its slogan “A  
3 sprinkle a day keeps odor away”, and through advertisements such as “Your body perspires in more  
4 places than just under your arms. Use SHOWER to SHOWER to feel dry, fresh, and comfortable  
5 throughout the day.” And “SHOWER to SHOWER can be used all over your body.”

6           16. In 1971, the first study was conducted that suggested an association between talc and  
7 ovarian cancer. This study was conducted by Dr. W.J. Henderson and others in Cardiff, Wales.

8           17. In 1982, the first epidemiologic study was performed on talc powder use in the  
9 female genital area. This study was conducted by Dr. Daniel Cramer and others. This study found a  
10 92% increased risk in ovarian cancer with women who reported genital talc use. Shortly after this  
11 study was published, Dr. Bruce Semple of Johnson & Johnson came and visited Dr. Cramer about  
12 his study. Dr. Cramer advised Dr. Semple that Johnson & Johnson should place a warning on its  
13 talcum powders about the ovarian cancer risks so that women can make an informed decision about  
14 their health.

15           18. Since 1982, there have been approximately twenty-two (22) additional  
16 epidemiologic studies providing data regarding the association of talc and ovarian cancer. Nearly  
17 all of these studies have reported an elevated risk for ovarian cancer associated with genital talc use  
18 in women.

19           a. In 1983, a case-control study found a 150% increased risk of ovarian cancer  
20 for women who use talcum powder in the genital area. Hartge, P., *et al.* Talc and Ovarian Cancer.  
21 *JAMA*. 1983; 250(14):1844.

22           b. In 1988, a case control study of 188 women diagnosed with epithelial  
23 ovarian cancer and 539 control women found that 52% of the cancer patients habitually used  
24 talcum powder on the genital area before their cancer diagnosis. The study showed a 50% increase  
25 in risk of ovarian cancer in women that used talcum powder on their genital area and a positive  
26 dose-response relationship. Whittemore AS, *et al.* Personal and environmental characteristics  
27 related to epithelial ovarian cancer. II. Exposures to talcum powder, tobacco, alcohol, and coffee.  
28 *Am. J. Epidemiol.* 1988 Dec; 128(6):1228-40.

1 c. A 1989 study looked at 235 women diagnosed with epithelial ovarian cancer  
2 and 451 controls, and found a 29% increased risk in ovarian cancer with women who reported  
3 genital talcum powder use more than once each week. Booth, M., *et al.* Risk factors for ovarian  
4 cancer: a case-control study. *Br J Cancer*. 1989 Oct; 60(4):592-8.

5 d. In 1992, a case-control study found a statistically significant 80% increased  
6 risk of ovarian cancer in women with more than 10,000 lifetime perineal applications of talc,  
7 demonstrating a positive dose-response relationship. Harlow BL, *et al.* Perineal exposure to talc and  
8 ovarian cancer risk. *Obstet Gynecol*. 1992 Jul; 80(1):19-26.

9 e. Another 1992 case-control study reported a 70% increased risk from genital  
10 talc use and a 379% increased risk of ovarian cancer of women who used talc on sanitary napkins  
11 in their genital area. Rosenblatt, K.A. *et al.* Mineral fiber exposure and the development of ovarian  
12 cancer. *Gynecol Oncol*. 1992 Apr; 45(1):20-5.

13 f. In 1995, the largest study of its kind to date found a statistically significant  
14 27% increased risk in ovarian cancer for women who regularly use talc in the abdominal or perineal  
15 area. Purdie, D., *et al.* Reproductive and other factors and risk of epithelial ovarian cancer: An  
16 Australian case-control study. Survey of Women's Health Study Group. *Int J Cancer*. 1995 Sep 15;  
17 62(6):678-84.

18 g. In 1996, a case-control study found a statistically significant 97% increased  
19 risk of ovarian cancer in women who used what they described as a "moderate" or higher use of  
20 talc-based powders in their genital area. *See* Shushan, A., *et al.* Human menopausal gonadotropin  
21 and the risk of epithelial ovarian cancer. *Fertil. Steril*. 1996 Jan; 65(1):13-8.

22 h. In 1997, a case control study of 313 women with ovarian cancer and 422  
23 without this disease found that the women with cancer were more likely to have applied talcum  
24 powder to their external genitalia area. Women using these products had a statistically significant  
25 50% to 90% higher risk of developing ovarian cancer. Cook, LS, *et al.* Perineal powder exposure  
26 and the risk of ovarian cancer. *Am. J Epidemiol*. 1997 Mar 1; 145(5):459-65.

27 i. In 1997, a case-control study involving over 1,000 women found a  
28 statistically significant increased risk of 42% for ovarian cancer for women who applied talc via

1 sanitary napkins to their perineal area. Chang, S, *et al.* Perineal talc exposure and risk of ovarian  
2 carcinoma. *Cancer*. 1997 Jun 15; 79(12):2396-401.

3 j. In 1998, a case-control study found a 149% increased risk of ovarian cancer  
4 in women who used talc-based powders on their perineal area. Godard, B., *et al.* Risk factors for  
5 familial and sporadic ovarian cancer among French Canadians: a case-control study. *Am J Obstet*  
6 *Gynecol.* 1998 Aug; 179(2):403-10.

7 k. Dr. Daniel Cramer conducted another case-control study in 1999, observing  
8 563 women newly diagnosed with epithelial ovarian cancer and 523 women in a control. The study  
9 found a statistically significant 60% increased risk of ovarian cancer in women that used talc-based  
10 body powders on their perineal area and an 80% increase in risk for women with over 10,000  
11 lifetime applications. Cramer, DW, *et al.* Genital talc exposure and risk of ovarian cancer. *Int J*  
12 *Cancer*. 1999 May 5; 81(3):351-56.

13 l. In 2000, a case-control study of over 2,000 women found a statistically  
14 significant 50% increased risk of ovarian cancer from genital talc use in women. Ness, RB, *et al.*  
15 Factors related to inflammation of the ovarian epithelium and risk of ovarian cancer. *Epidemiology*.  
16 2000 Mar; 11(2):111-7.

17 m. In 2004, a case-control study of nearly 1,400 women from 22 counties in  
18 Central California found a statistically significant 37% increased risk of epithelial ovarian cancer  
19 from women's genital talc use, and a 77% increased risk of serous invasive ovarian cancer from  
20 women's genital talc use. Importantly, this study also examined at women's use of cornstarch  
21 powders as an alternative to talc, and found no increased risk in ovarian cancer in women in the  
22 cornstarch group, further supporting the causal connection between genital talc use and ovarian  
23 cancer. Mills, PK, *et al.* Perineal talc exposure and epithelial ovarian cancer risk in the Central  
24 Valley of California. *Int J Cancer*. 2004 Nov 10; 112(3):458-64.

25 n. In 2008, a combined study of over 3,000 women from a New England-based  
26 case-control study found a general 36% statistically significant increased risk of epithelial ovarian  
27 cancer from genital talc use and a 60% increased risk of the serous invasive ovarian cancer subtype.  
28 The study also found a strong dose-response relationship between the cumulative talc exposure and

1 incidence of ovarian cancer, adding further support to the causal relationship. Gates, MA, *et al.*  
2 Talc Use, Variants of the GSTM1, GSTT1, and NAT2 Genes, and Risk of Epithelial Ovarian  
3 Cancer. *Cancer Epidemiol Biomarkers Prev.* 2008 Sep; 17(9):2436-44.

4 o. A 2009 case-control study of over 1,200 women found the risk of ovarian  
5 cancer increased significantly with increasing frequency and duration of talc use, with an overall  
6 statistically significant 53% increased risk of ovarian cancer from genital talc use. That increased  
7 risk rose dramatically, to 108%, in women with the longest duration and most frequent talc use.  
8 Wu, AH, *et al.* Markers of inflammation and risk of ovarian cancer in Los Angeles County. *Int. J*  
9 *Cancer.* 2009 Mar 15; 124(6):1409-15.

10 p. In 2011, another case-control study of over 2,000 women found a 27%  
11 increased risk of ovarian cancer from genital talc use. Rosenblatt, KA, *et al.* Genital powder  
12 exposure and the risk of epithelial ovarian cancer. *Cancer Causes Control.* 2011 May; 22(5):737-  
13 42.

14 q. In June of 2013, a pooled analysis of over 18,000 women in eight case-  
15 control studies found a 20% to 30% increased risk of women developing epithelial ovarian cancer  
16 from genital powder use. The study concluded by stating, “Because there are few modifiable risk  
17 factors for ovarian cancer, avoidance of genital powders may be a possible strategy to reduce  
18 ovarian cancer incidence.” Terry, KL, *et al.* Genital powder use and risk of ovarian cancer: a pooled  
19 analysis of 8,525 cases and 9,859 controls. *Cancer Prev Res (Phila).* 2013 Aug; 6(8):811-21.

20 19. Researchers have also examined the link between endometrial cancer, a form of  
21 uterine cancer, and the application of talcum powder to the perineal area.

22 20. In 2010, one such study analyzed data from a 1976 cohort study of over 66,000  
23 women, and found a statistically significant 21% increased risk of endometrial (uterine) cancer in  
24 postmenopausal women who had ever applied talcum powder in the perineal area. This risk rose to  
25 24% for postmenopausal women who applied talc in the perineal area “regularly,” defined as at  
26 least once a week. Karageorgi S., *et al.* (2010) Perineal use of talcum powder and endometrial  
27 cancer risk. *Cancer Epidemiol Biomarkers Prev.* 2010 May; 19:1269–1275.

28

1           21.     In 1993, the United States National Toxicology Program published a study on the  
2 toxicity of non-asbestiform talc and found clear evidence of carcinogenic activity. Talc was found  
3 to be a carcinogen, with or without the presence of asbestos-like fibers.

4           22.     In response to the United States National Toxicology Program’s study, the Cosmetic  
5 Toiletry and Fragrance Association (CTFA) formed the Talc Interested Party Task Force (TIPTF).  
6 Johnson & Johnson, Inc., and Johnson & Johnson Consumer Companies, Inc. were members of the  
7 CTFA. The stated purpose of the TIPTF was to pool financial resources of these companies in an  
8 effort to collectively defend talc use at all costs and to prevent regulation of any type over this  
9 industry. The TIPTF hired scientists to perform biased research regarding the safety of talc,  
10 members of the TIPTF edited scientific reports of the scientists hired by this group prior to the  
11 submission of these scientific reports to governmental agencies, members of the TIPTF knowingly  
12 released false information about the safety of talc to the consuming public, and used political and  
13 economic influence on regulatory bodies regarding talc. All of these activities have been well  
14 coordinated and planned by these companies and organizations over the past four (4) decades in an  
15 effort to prevent regulation of talc and to create confusion to the consuming public about the true  
16 hazards of talc relative to cancer.

17           23.     On November 10, 1994, the Cancer Prevention Coalition mailed a letter to then  
18 Johnson & Johnson C.E.O, Ralph Larson, informing his company that studies as far back as 1960’s  
19 “. . . show[ ] conclusively that the frequent use of talcum powder in the genital area pose[ ] a  
20 serious health risk of ovarian cancer.” The letter cited a recent study by Dr. Bernard Harlow from  
21 Harvard Medical School confirming this fact and quoted a portion of the study where Dr. Harlow  
22 and his colleagues discouraged the use of talc in the female genital area. The letter further stated  
23 that 14,000 women per year die from ovarian cancer and that this type of cancer is very difficult to  
24 detect and has a low survival rate. The letter concluded by requesting that Johnson & Johnson  
25 withdraw talc products from the market because of the alternative of cornstarch powders, or at a  
26 minimum, place warning information on its talc-based body powders about ovarian cancer risk they  
27 pose.

28



1           24.     In 1996, the condom industry stopped dusting condoms with talc due to the growing  
2 health concerns.

3           25.     In February of 2006, the International Association for the Research of Cancer  
4 (IARC) part of the World Health Organization published a paper whereby they classified perineal  
5 use of talc based body powder as a “Group 2B” human carcinogen. IARC which is universally  
6 accepted as the international authority on cancer issues, concluded that studies from around the  
7 world consistently found an increased risk of ovarian cancer in women from perineal use of talc.  
8 IARC found that between 16-52% of women in the world were using talc to dust their perineum  
9 and found an increased risk of ovarian cancer in women talc users ranging from 30-60%. IARC  
10 concluded with this “Evaluation”: “There is limited evidence in humans for the carcinogenicity of  
11 perineal use of talc-based body powder.” By definition “Limited evidence of carcinogenicity”  
12 means “a positive association has been observed between exposure to the agent and cancer for  
13 which a causal interpretation is considered by the Working Group to be credible, but chance, bias or  
14 confounding could not be ruled out with reasonable confidence.”

15           26.     In approximately 2006, the Canadian government under The Hazardous Products  
16 Act and associated Controlled Products Regulations classified talc as a “D2A,” “very toxic,” 51  
17 “cancer causing” substance under its Workplace Hazardous Materials Information System  
18 (WHMIS). Asbestos is also classified as “D2A”.

19           27.     In 2006, Imerys Talc began placing a warning on the Material Safety Data Sheets  
20 (MSDS) it provided to the Johnson & Johnson Defendants regarding the talc it sold to them to be  
21 used in the Products. These MSDSs not only provided the warning information about the IARC  
22 classification but also included warning information regarding “States Rights to Know” and  
23 warning information about the Canadian Government’s “D2A” classification of talc as well.

24           28.     Defendants had a duty to know and warn about the hazards associated with the use  
25 of the Products.

26           29.     Defendants failed to inform customers and end users of the Products of a known  
27 catastrophic health hazard associated with the use of the Products.

28

1 30. In addition, Defendants procured and disseminated false, misleading, and biased  
2 information regarding the safety of the Products to the public and used influence over governmental  
3 and regulatory bodies regarding talc.

4 **B. Plaintiff's Use of the Products**

5 31. Plaintiff was born in 1975, and is a resident of Contra Costa County, California.

6 32. When Plaintiff was an infant, her mother used Shower to Shower, and applied J&J  
7 Baby Powder to Plaintiff. As she grew up, Plaintiff liked the smell and feel of the Products, and  
8 used them continuously until she entered the military at the Great Lakes Naval Station.

9 33. Plaintiff continued to use the Products following her discharge from the Navy.

10 34. There was never any indication, on the Products packaging or otherwise, that this  
11 normal use could and would cause her to develop uterine cancer.

12 35. Plaintiff was diagnosed with uterine cancer in 2006.

13 36. In 2007, Plaintiff underwent surgical removal of her uterus, ovaries and fallopian  
14 tubes.

15 37. Plaintiff's uterine cancer has been in remission since 2007.

16 **COUNT ONE - STRICT LIABILITY**  
17 **(FAILURE TO WARN)**

18 38. Plaintiff incorporates by reference each of the preceding paragraphs as if fully set  
19 forth herein.

20 39. At all pertinent times, the Johnson & Johnson Defendants were manufacturing,  
21 marketing, testing, promoting, selling and/or distributing the Products in the regular course of  
22 business.

23 40. At all pertinent times, Plaintiff used the Products to powder her perineal area, which  
24 is a reasonably foreseeable use.

25 41. At all pertinent times, Defendants in this action knew or should have known that the  
26 use of talcum powder based products in the perineal area significantly increases the risk of cancer,  
27 including, but not limited to, ovarian and uterine cancer, based upon scientific knowledge dating  
28 back for decades.

1           42. At all pertinent times, including the time of sale and consumption, the Products,  
2 when put to the aforementioned reasonably foreseeable use, were in an unreasonably dangerous and  
3 defective condition because they failed to contain adequate and proper warnings and/or instructions  
4 regarding the increased risk of cancer, including, but not limited to, ovarian and uterine cancer,  
5 associated with the use of the Products by women to powder their perineal area. Defendants  
6 themselves failed to properly and adequately warn and instruct Plaintiff as to the risks and benefits  
7 of the Products given her need for this information.

8           43. Had Plaintiff received a warning that the use of the Products would significantly  
9 increase her risk of developing cancer, she would not have used them. As a proximate result of  
10 Defendants' design, manufacture, marketing, sale, and distribution of the Products, Plaintiff was  
11 injured catastrophically, and was caused severe pain, suffering, disability, impairment, loss of  
12 enjoyment of life, loss of care, comfort, and economic damages.

13           44. The development of uterine cancer by Plaintiff was the direct and proximate result of  
14 the unreasonably dangerous and defective condition of the Products at the time of sale and  
15 consumption, including their lack of warnings; Plaintiff suffered injuries and damages including,  
16 but not limited to, physical and mental pain and suffering, and medical expenses.

17           45. Defendants' products were defective because they failed to contain warnings and/or  
18 instructions, and breached express warranties and/or failed to conform to express factual  
19 representations upon which Plaintiff justifiably relied in electing to use the Products. The defect or  
20 defects made the Products unreasonably dangerous to persons, such as Plaintiff, who could  
21 reasonably be expected to use and rely upon such products. As a result, the defect or defects were a  
22 producing cause of Plaintiff's injuries and damages.

23           46. Defendants' products failed to contain, and continue to this day not to contain,  
24 adequate warnings and/or instructions regarding the increased risk of cancer, including, but not  
25 limited to, ovarian and uterine cancer, with the use of their products by women. Defendants  
26 continue to market, advertise, and expressly represent to the general public that it is safe for women  
27 to use their product regardless of application. These Defendants continue with these marketing and  
28

1 advertising campaigns despite having scientific knowledge that dates back to the 1960's that their  
2 products increase the risk of cancer in women when used in the perineal area.

3 47. Plaintiff sustained the following damages as a foreseeable, direct, and proximate  
4 result of Defendants' acts and/or omissions:

- 5 a. Economic losses including medical care and lost earnings; and  
6 b. Noneconomic losses including physical and mental pain and suffering,  
7 emotional distress, inconvenience, loss of enjoyment and impairment of quality of life, past and  
8 future.

9 **COUNT TWO – STRICT LIABILITY**  
10 **(DESIGN AND/OR MANUFACTURING DEFECT)**

11 48. Plaintiff incorporates by reference each of the preceding paragraphs as if fully set  
12 forth herein.

13 49. Defendants engaged in the design, development, manufacture, marketing, sale, and  
14 distribution of the Products in a defective and unreasonably dangerous condition to consumers,  
15 including Plaintiff.

16 50. Defendants caused the Products to enter the stream of commerce and to be sold  
17 through various retailers, where Plaintiff purchased the Products.

18 51. The Products were expected to, and did, reach consumers, including Plaintiff,  
19 without change in the condition in which it was manufactured and sold by Defendants and/or  
20 otherwise released into the stream of commerce.

21 52. Plaintiff used the Products in a manner normally intended, recommended, promoted,  
22 and marketed by Defendants.

23 53. Products failed to perform safely when used by Plaintiff in a reasonably foreseeable  
24 manner, specifically increasing her of developing uterine cancer.

25 54. The propensity of talc fibers to translocate into the female reproductive system,  
26 including, but not limited to, the ovaries and endometrial lining of the uterus, thereby substantially  
27 increasing the risk of cancer, including, but not limited to, ovarian and uterine cancer, renders the  
28 Products unreasonably dangerous when used in the manner it was intended and to an extent beyond  
that would be contemplated by the ordinary consumer.

1 55. Importantly, the Products are an inessential cosmetic product that do not treat or cure  
2 any serious disease. Further, safer alternatives, including corn-starch based powders, have been  
3 readily available for decades.

4 56. Defendants have known, or should have known, that the Products are unreasonably  
5 dangerous when used by a woman in her perineal area but have continued to design, manufacture,  
6 sell, distribute, market, promote, and supply the Products so as to maximize sales and profits at the  
7 expense of public health and safety in conscious disregard of the foreseeable harm to the consuming  
8 public, including Plaintiff.

9 57. As a direct and proximate result of Defendants' conduct, including actions,  
10 omissions, and misrepresentations, Plaintiff sustained the following damages:

- 11 a. Economic losses including medical care and lost earnings; and
- 12 b. Noneconomic losses including physical and mental pain and suffering, emotional  
13 distress, inconvenience, loss of enjoyment and impairment of quality of life, past and future.

14 **COUNT THREE-NEGLIGENCE**

15 58. Plaintiff incorporates by reference each of the preceding paragraphs as if fully set  
16 forth herein.

17 59. The Johnson & Johnson Defendants were negligent in marketing, designing,  
18 manufacturing, producing, supplying, inspecting, testing, selling and/or distributing the Products in  
19 one or more of the following respects:

- 20 • In failing to warn Plaintiff of the hazards associated with the use of the Products;
- 21 • In failing to properly test their products to determine adequacy and effectiveness or  
22 safety measures, if any, prior to releasing the Products for consumer use;
- 23 • In failing to properly test their products to determine the increased risk of ovarian  
24 cancer during the normal and/or intended use of the Products;
- 25 • In failing to inform ultimate users, such as Plaintiff as to the safe and proper  
26 methods of handling and using the Products;
- 27 • In failing to remove the Products from the market when Defendants knew or should  
28 have known the Products were defective;

- 1 • In failing to instruct the ultimate users, such as Plaintiff, as to the methods for
- 2 reducing the type of exposure to the Products which caused increased risk of cancer,
- 3 including, but not limited to, ovarian and uterine cancer;
- 4 • In failing to inform the public in general and Plaintiff in particular of the known
- 5 dangers of using the Products for dusting the perineum;
- 6 • In failing to advise users how to prevent or reduce exposure that caused increased
- 7 risk for cancer, including, but not limited to, ovarian and uterine cancer;
- 8 • In marketing and labeling the Products as safe for all uses despite knowledge to the
- 9 contrary.
- 10 • In failing to act like a reasonably prudent company under similar circumstances.

11 Each and all of these acts and omissions, taken singularly or in combination, were a  
12 proximate cause of the injuries and damages sustained by Plaintiff.

13 60. At all pertinent times, the Johnson & Johnson Defendants knew or should have  
14 known that the Products were unreasonably dangerous and defective when put to their reasonably  
15 anticipated use.

16 61. Plaintiff sustained the following damages as a foreseeable, direct, and proximate  
17 result of Defendants' acts and/or omissions:

- 18 a. Economic losses including medical care and lost earnings; and
- 19 b. Noneconomic losses including physical and mental pain and suffering,
- 20 emotional distress, inconvenience, and loss of enjoyment and impairment of quality of life, past and
- 21 future.

22 **COUNT FOUR- BREACH OF EXPRESS WARRANTY**

23 62. Plaintiff incorporates by reference each of the preceding paragraphs as if fully set  
24 forth herein.

25 63. The Johnson & Johnson Defendants expressly warranted, through direct-to-  
26 consumer marketing, advertisements, and labels, that the Products were safe and effective for  
27 reasonably anticipated uses, including use by women in the perineal area.

28

1           64. The Products did not conform to these express representations because they cause  
2 serious injury when used by women in the perineal area in the form of cancer, including, but not  
3 limited to, ovarian and uterine cancer.

4           65. Plaintiff sustained the following damages as a foreseeable, direct, and proximate  
5 result of Defendants' acts and/or omissions:

- 6           a. Economic losses including medical care and lost earnings; and
- 7           b. Noneconomic losses including physical and mental pain and suffering,  
8 emotional distress, inconvenience, loss of enjoyment and impairment of quality of life, past and  
9 future.

10                                   **COUNT FIVE – BREACH OF IMPLIED WARRANTIES**

11           66. Plaintiff incorporates by reference each of the preceding paragraphs as if fully set  
12 forth herein.

13           67. At the time the Defendants manufactured, marketed, labeled, promoted, distributed  
14 and/or sold the Products, the Johnson & Johnson Defendants knew of the uses for which the  
15 Products were intended, including use by women in the perineal area, and impliedly warranted the  
16 Products to be of merchantable quality and safe for such use.

17           68. Defendants breached their implied warranties of the Products sold to Plaintiff  
18 because they were not fit for their common, ordinary and intended uses, including use by women in  
19 the perineal area.

20           69. Plaintiff sustained the following damages as a foreseeable, direct, and proximate  
21 result of Defendants' acts and/or omissions:

- 22           a. Economic losses including medical care and lost earnings; and
- 23           b. Noneconomic losses including physical and mental pain and suffering,  
24 emotional distress, inconvenience, loss of enjoyment and impairment of quality of life, past and  
25 future.

26                                   **COUNT SIX – PUNITIVE DAMAGES**

27           70. Plaintiff incorporates by reference each of the preceding paragraphs as if fully set  
28 forth herein.

1 71. Defendants have acted willfully, wantonly, with an evil motive, and recklessly in  
2 one or more of the following ways:

- 3 a. Defendants knew of the unreasonably high risk of cancer, including, but not limited  
4 to, ovarian and uterine cancer, posed by the Products before manufacturing,  
5 marketing, distributing and/or selling the Products, yet purposefully proceeded with  
6 such action;
- 7 b. Despite their knowledge of the high risk of cancer, including, but not limited to,  
8 ovarian and uterine cancer, associated with the Products, Defendants affirmatively  
9 minimized this risk through marketing and promotional efforts and product labeling;
- 10 c. Through the actions outlined above, Defendants expressed a reckless indifference to  
11 the safety of users of the Products, including Plaintiff. Defendants' conduct, as  
12 described herein, knowing the dangers and risks of the Products, yet concealing  
13 and/or omitting this information, in furtherance of their conspiracy and concerted  
14 action was outrageous because of Defendants' evil motive or a reckless indifference  
15 to the safety of users of the Products.

16 72. Plaintiff sustained the following damages as a foreseeable, direct, and proximate  
17 result of Defendants' acts and/or omissions:

- 18 a. Economic losses including medical care and lost earnings; and  
19 b. Noneconomic losses including physical and mental pain and suffering,  
20 emotional distress, inconvenience, loss of enjoyment and impairment of quality of life, past and  
21 future.

22 **COUNT SEVEN – NEGLIGENT MISREPRESENTATION**

23 73. Plaintiff incorporates by reference each of the preceding paragraphs as if fully set  
24 forth herein.

25 74. Defendants had a duty to accurately and truthfully represent to the medical and  
26 healthcare community, Plaintiff, and the public, that the Products had been tested and found to be  
27 safe and effective for use in the perineal area. The representations made by Defendants, in fact,  
28 were false.

75. Defendants failed to exercise ordinary care in the representations concerning the  
Products while they were involved in their manufacture, sale, testing, quality assurance, quality



1 control, and distribution in interstate commerce, because Defendants negligently misrepresented the  
2 Products' high risk of unreasonable, dangerous, adverse side effects.

3 76. Defendants breached their duty in representing that the Products have no serious side  
4 effects.

5 77. As a foreseeable, direct and proximate result of the negligent misrepresentation of  
6 Defendants as set forth herein, Defendants knew, and had reason to know, that the Products had  
7 been insufficiently tested, or had not been tested at all, and that they lacked adequate and accurate  
8 warnings, and that it created a high risk, and/or higher than acceptable risk, and/or higher than  
9 reported and represented risk, of adverse side effects, including, but not limited to, ovarian and  
10 uterine cancer.

11 78. Plaintiff sustained the following damages as a foreseeable, direct, and proximate  
12 result of Defendants' acts and/or omissions:

- 13 a. Economic losses including medical care and lost earnings; and  
14 b. Noneconomic losses including physical and mental pain and suffering,  
15 emotional distress, inconvenience, loss of enjoyment and impairment of quality of life, past and  
16 future.

17 **COUNT EIGHT – FRAUDULENT CONCEALMENT**

18 79. Plaintiff incorporates by reference each of the preceding paragraphs as if fully set  
19 forth herein.

20 80. Defendants owed consumers, including Plaintiff, a duty to fully and accurately  
21 disclose all material facts regarding the Products, not to conceal material defects related thereto, not  
22 to place these defective products into the stream of commerce, and to fully and accurately label  
23 product packaging. To the contrary, Defendants explicitly and/or implicitly represented that the  
24 Products were safe and effective.

25 81. Defendants actively and intentionally concealed and/or suppressed material facts, in  
26 whole or in part, to induce consumers, including Plaintiff, to purchase and use the Products and did  
27 so at her expense. Specifically:

- 28 a. Defendants have been aware of the positive association between feminine

1 talc use and cancer demonstrated by epidemiology studies since at least 1982 and more than a  
2 dozen such published studies, including meta- analyses, have been published demonstrating similar  
3 results;

4 b. Defendants have been aware, for decades, of the propensity for talc particles  
5 to translocate from the perineum through the vaginal tract into the ovaries;

6 c. IARC, the recognized world authority of agent carcinogenicity, has  
7 determined that there is a credible causal connection between feminine talc use and ovarian cancer;  
8 and

9 d. Johnson & Johnson's own paid consultant, Dr. Alfred Wehner, advised the  
10 company on multiple occasions, by at least 1997, that Johnson & Johnson's denial of a positive  
11 association between feminine talc use and ovarian cancer was "technically and factually incorrect."

12 e. Recent studies have established a statistically significant correlation between  
13 talcum powder use in the perineal area and uterine cancer.

14 82. Defendants made the misrepresentations and/or omissions for the purpose of  
15 deceiving and defrauding Plaintiff and with the intention of having her act and rely on such  
16 misrepresentations and/or omissions.

17 83. Defendants knew that their concealments, misrepresentations and/or omissions were  
18 material, and that they were false, incomplete, misleading, deceptive, and deceitful when they were  
19 made. Alternatively, Defendants concealed information, and/or made the representations with such  
20 reckless disregard for the truth that knowledge of the falsity can be imputed to them.

21 84. Defendants profited, significantly, from their unethical and illegal conduct that  
22 caused Plaintiff to purchase and habitually use a dangerous and defective product.

23 85. Defendants' actions, and Plaintiff's justifiable reliance thereon, were substantial  
24 contributing factors in causing injury and incurrence of substantial damages.

25 86. Plaintiff sustained the following damages as a foreseeable, direct, and proximate  
26 result of Defendants' acts and/or omissions:

27 a. Economic losses including medical care and lost earnings; and

28 b. Noneconomic losses including physical and mental pain and suffering,

1 emotional distress, inconvenience, loss of enjoyment and impairment of quality of life, past and  
2 future.

3 **COUNT NINE – FRAUD**  
4 **(INTENTIONAL MISREPRESENTATION)**

5 87. Plaintiff incorporates by reference each of the preceding paragraphs as if fully set  
6 forth herein.

7 88. Defendants, who engaged in the development, manufacture, marketing, sale and  
8 distribution of personal hygiene products, including the Products, owed a duty to provide accurate  
9 and complete information regarding said products.

10 89. Defendants fraudulently misrepresented the use of the Products as safe and  
11 effective, specifically:

12 a. Johnson & Johnson’s website calls it a “misconception” that talc in baby  
13 powder can be “absorbed into the body”;

14 b. Johnson & Johnson print advertisements directed at adult women asserted  
15 that, because Johnson & Johnson Baby Powder is used on babies, women can “trust” that Johnson  
16 & Johnson will take “just as much care” of their skin;

17 c. Misleading consumers in advertisements that the talc in Johnson & Johnson  
18 Baby Powder is safe because it comes from “nature” and is “pure”;

19 d. Johnson & Johnson, on its website, claims that “30 years of research by  
20 independent scientists, review boards and global authorities [] have concluded that talc can be used  
21 safely in personal care products,” failing to mention the dozens of studies demonstrating a  
22 relationship between feminine talc use and ovarian cancer, as well as the decision by IARC to label  
23 feminine talc powder use as “possibly carcinogenic”; and

24 e. On the Johnson & Johnson Baby Powder bottle, Defendants include a  
25 conspicuous warning to mothers to prevent babies from inhaling the powder and the inclusion of  
26 this lone warning implies to the consumer that Johnson & Johnson Baby Powder is safe in all other  
27 manners of use.

28 90. Defendants knew that these misrepresentations and/or omissions were material, and  
that they were false, incomplete, misleading, deceptive and deceitful when they were made.

1 91. Defendants made the misrepresentations and/or omissions for the purpose of  
2 deceiving and defrauding consumers, including Plaintiff, with the intention of having them act and  
3 rely on such misrepresentations and/or omissions.

4 92. Plaintiff relied, with reasonable justification, on the misrepresentations by  
5 Defendants, which induced her to purchase and use the Products on a regular basis for decades.

6 93. Defendants profited, significantly, from their unethical and illegal conduct that  
7 fraudulently induced Plaintiff, and millions of other consumers, to purchase a dangerous and  
8 defective product.

9 94. Defendants' actions, and Plaintiff's justifiable reliance thereon, were substantial  
10 contributing factors in causing injury and incurrence of substantial damages.

11 95. As a foreseeable, direct, and proximate result of the aforementioned fraudulent  
12 misrepresentations by Defendants, Plaintiff sustained the following damages:

- 13 a. Economic losses including medical care and lost earnings; and  
14 b. Noneconomic losses including physical and mental pain and suffering,  
15 emotional distress, inconvenience, loss of enjoyment and impairment of quality of life, past and  
16 future.

17 **COUNT TEN – VIOLATION OF THE UCL**

18 96. Plaintiff incorporates by reference each of the preceding paragraphs as if fully set  
19 forth herein.

20 97. California's UCL prohibits any "unlawful, unfair, or fraudulent" business practice.  
21 Cal. Bus. & Prof. Code. § 17200. Defendants' misrepresentations and omissions described herein  
22 are "unlawful, unfair and fraudulent" under California law.

23 98. Plaintiff purchased and used the Johnson & Johnson Defendants' Products primarily  
24 for personal use and thereby suffered ascertainable losses as a result of Defendants' actions in  
25 violation of the UCL.

26 99. Had Defendants not engaged in the deceptive conduct described herein, Plaintiff  
27 would not have purchased and/or paid for Defendants' Products, and would not have incurred  
28 related injuries and damages.

1           100. Defendants engaged in wrongful conduct while at the same time obtaining, under  
2 false pretenses, monetary gain from Plaintiff for the Products that would not have been paid had  
3 Defendants not engaged in fraudulent conduct.

4           101. Defendants engaged in fraudulent methods of competition and deceptive acts or  
5 practices that were proscribed by law, including the following:

6                 a. Representing that goods or services have characteristics, ingredients, uses,  
7 benefits, or quantities that they do not have;

8                 b. Advertising goods or services with the intent not to sell them as advertised; and

9                 c. Engaging in fraudulent conduct that creates a likelihood of confusion or  
10 misunderstanding.

11           102. Defendants intended for Plaintiff to rely on their representations and  
12 advertisements regarding the Products in order to achieve monetary gain from Plaintiff through her  
13 purchase of the Products.

14           103. Plaintiff was injured by the cumulative and indivisible nature of Defendants'  
15 conduct. The cumulative effect of Defendants' conduct directed at Plaintiff and other consumers  
16 was to create demand for and sell the Products. Each aspect of Defendants' conduct combined to  
17 artificially create sales of the Products.

18           104. Defendants have a statutory duty to refrain from unfair or deceptive acts or trade  
19 practices in the design, labeling, development, manufacture, promotion, and sale of the Products.

20           105. Had Defendants not engaged in the deceptive conduct described above, Plaintiff  
21 would not have purchased and/or paid for the product, and would not have incurred related injuries  
22 and damages.

23           106. Defendants' intentional, deceptive, unconscionable, and fraudulent representations  
24 and material omissions to Plaintiff, physicians, and consumers, constituted unfair and deceptive acts  
25 and trade practices in violation of Cal. Bus. & Prof. Code. § 17200.

26           107. Defendants' actions, as complained of herein, constitute unfair competition or unfair,  
27 unconscionable, deceptive or fraudulent acts, or trade practices in violation of Cal. Bus. & Prof.  
28 Code. § 17200.

1 108. Defendants have engaged in unfair competition or unfair or deceptive acts or trade  
2 practices, or have made false representations in violation of Cal. Bus. & Prof. Code. § 17200.

3 109. Defendants are the suppliers, manufacturers, advertisers, and sellers of the Products,  
4 and are subject to liability under Cal. Bus. & Prof. Code. § 17200 for unfair, deceptive, fraudulent  
5 and unconscionable consumer sales practices.

6 110. Defendants violated Cal. Bus. & Prof. Code. § 17200, by knowingly and falsely  
7 representing that Defendants' Products were fit to be used for the purpose for which they were  
8 intended, when in fact the Products were and are defective and dangerous, and by other acts alleged  
9 herein. These representations were made in marketing and promotional materials.

10 111. Defendants had actual knowledge of the defective and dangerous condition of  
11 Defendants' Products, and failed to take any action to cure such defective and dangerous  
12 conditions.

13 112. Plaintiff relied upon Defendants' misrepresentations and omissions in determining  
14 which Products to use.

15 113. Defendants' deceptive, unconscionable or fraudulent representations and material  
16 omissions to Plaintiff and other consumers constituted deceptive acts and practices.

17 114. By reason of the unlawful acts engaged in by Defendants, and as a direct and  
18 proximate result thereof, Plaintiff suffered ascertainable losses and damages.

19 115. As a direct and proximate result of Defendants' violations of Cal. Bus. & Prof.  
20 Code. § 17200, Plaintiff sustained the following damages:

- 21 a. Economic losses including medical care and lost earnings; and  
22 b. Noneconomic losses including physical and mental pain and suffering,  
23 emotional distress, inconvenience, loss of enjoyment and impairment of quality of life, past and  
24 future.

25 **COUNT ELEVEN – RESTITUTION OR DISGORGEMENT BASED ON UNJUST**  
26 **ENRICHMENT**

27 116. Plaintiff incorporates by reference each of the preceding paragraphs as if fully set  
28 forth herein.

1 117. As a result of the Johnson & Johnson Defendants' unlawful, fraudulent and  
2 misleading labeling, advertising, marketing and sales of the Products described herein, Defendants  
3 were unjustly enriched at the expense of Plaintiff.

4 118. Defendants sold their Products to Plaintiff as described herein, and profited  
5 therefrom. It would be against equity and good conscience to permit Defendants to retain the ill-  
6 gotten benefits Defendants received from Plaintiff, in light of the fact that the Products were not  
7 what Defendants purported them to be. Thus, it would be unjust and inequitable for Defendants to  
8 retain the benefit without restitution or disgorgement to Plaintiff of monies paid to Defendants for  
9 the Products.

10 **COUNT TWELVE - CONSUMER LEGAL REMEDIES ACT**

11 119. Plaintiff incorporates by reference each of the preceding paragraphs as if fully set  
12 forth herein.

13 120. This cause of action is brought under the Consumers Legal Remedies Act, California  
14 Civil Code §§ 1750, et seq.

15 121. Plaintiff presently seeks only injunctive relief under this cause of action. Plaintiff  
16 will amend this cause of action to seek damages after giving the notice required by Cal. Civ. Code §  
17 1782.

18 122. Plaintiff is a "consumer" within the meaning of Civil Code § 1761(d).

19 123. Defendants' sales of their Products constitute "transactions" within the meaning of  
20 Civil Code § 1761(e). The Products purchased by Plaintiff constitute "goods" under Civil Code §  
21 1761(a).

22 124. As described above, Defendants' representations to Plaintiff were false, in violation  
23 of the CLRA. Defendants' conduct violated, among others, (1) Civil Code § 1770(a)(5), which  
24 prohibits "[r]epresenting that goods or services have sponsorship, approval, characteristics,  
25 ingredients, uses, benefits, or quantities which they do not have or that a person has a sponsorship,  
26 approval, status, affiliation, or connection which he or she does not have"; (2) Civil Code §  
27 1770(a)(7), which prohibits "[r]epresenting that goods or services are of a particular standard,  
28 quality, or grade, or that goods are of a particular style or model, if they are of another"; and (3)

1 Civil Code § 1770(a)(9), which prohibits “[a]dvertising goods or services with intent not to sell  
2 them as advertised.”

3 125. The violations of the CLRA by Defendants were willful, oppressive, and fraudulent.

4 126. Pursuant to Cal. Civ. Code § 1782(a)(2), Plaintiff is entitled to an order enjoining the  
5 above-described acts and practices.

6 **COUNT 13 – FALSE ADVERTISING LAW**

7 127. Plaintiff incorporates by reference each of the preceding paragraphs as if fully set  
8 forth herein.

9 128. This cause of action is brought under California’s False Advertising Law, California  
10 Business & Professions Code §§ 17500, et seq.

11 129. The FAL prohibits the dissemination of any advertising which is untrue or  
12 misleading, and which is known, or which by the exercise of reasonable care should be known, to  
13 be untrue or misleading. Cal. Bus. & Prof. Code § 17500.

14 130. The Johnson & Johnson Defendants engaged in a scheme of offering the Products  
15 described herein for sale to Plaintiff by way of advertising, product packaging and labeling, and  
16 other promotional materials. Defendants misrepresented the true contents and nature of Defendants’  
17 Products.

18 131. As explained herein, Defendants advertised, and continue to advertise, its Products  
19 in a manner that was, and is, untrue and misleading.

20 132. Defendants knew or should have known that their advertisements were and are  
21 misleading or likely to mislead for the reasons set forth above.

22 133. Defendants’ advertisements and inducements were made within California and come  
23 within the definition of advertising as contained in Business and Professions Code §17500, et seq.

24 134. Defendants’ Product packaging and labeling, and promotional materials, were  
25 intended as inducements to purchase Defendants’ Products, and are statements disseminated by  
26 Defendants to Plaintiff.

27 135. Defendants’ advertisements induced Plaintiff to purchase Defendants’ Products, as  
28 described herein.



1           136. Plaintiff suffered injuries in fact and losses of money or property as a result of  
2 Defendants' acts and practices, which violate §§ 17500, et seq.

3   **TOLLING OF STATUTE OF LIMITATIONS**

4           137. Plaintiff incorporates by reference each of the preceding paragraphs as if fully set  
5 forth herein.

6           138. Plaintiff suffered an illness that had a latency period and did not arise until many  
7 years after exposure. Plaintiff was not aware at the time of her diagnosis that her uterine cancer was  
8 caused by her use of the Defendants' Products. Consequently, the discovery rule applies to this case  
9 and the statute of limitations has been tolled until the day that Plaintiff knew or had reason to know  
10 that her uterine cancer was linked to her use of Defendants' Products.

11           139. Furthermore, the running of any statute of limitations has been equitably tolled by  
12 reason of Defendants' fraudulent concealment and conduct. Through their affirmative  
13 misrepresentations and omissions, Defendants actively concealed from Plaintiff the true risks  
14 associated with the Products.

15           140. As a result of Defendants' actions, Plaintiff and her prescribing physicians were  
16 unaware, and could not reasonably know or have learned through reasonable diligence that she had  
17 been exposed to the risks alleged herein and that those risks were the direct and proximate result of  
18 Defendants' acts and omissions.

19           141. Furthermore, Defendants are estopped from relying on any statute of limitations  
20 because of their concealment of the truth, quality and nature of the Products. Defendants were  
21 under a duty to disclose the true character, quality and nature of the Products because this was non-  
22 public information over which the Defendants had and continue to have exclusive control, and  
23 because the Defendants knew that this information was not available to Plaintiff, her medical  
24 providers and/or her health facilities.

25           142. Defendants had the ability to and did spend enormous amounts of money in  
26 furtherance of their purpose of marketing and promoting a profitable product, notwithstanding the  
27 known or reasonably known risks. Plaintiff and medical professionals could not have afforded and  
28

1 could not have possibly conducted studies to determine the nature, extent and identity of related  
2 health risks, and were forced to rely on Defendants' representations.

3 **PRAYER FOR RELIEF**

4 WHEREFORE, Plaintiff demands judgment against Defendants on each of the above-  
5 referenced claims and causes of action, and as follows:

6 a. Awarding compensatory damages in excess of \$75,000, including, but not  
7 limited to pain, suffering, emotional distress, loss of enjoyment of life, and other non-economic  
8 damages in an amount to be determined at trial of this action;

9 b. Awarding economic damages in the form of medical expenses, out of pocket  
10 expenses, lost earnings, and other economic damages in an amount to be determined at trial of this  
11 action;

12 c. Punitive and/or exemplary damages for the wanton, willful, fraudulent,  
13 reckless acts of the Defendants who demonstrated a complete disregard and reckless indifference  
14 for the safety and welfare of the general public and Plaintiff in an amount sufficient to punish  
15 Defendants and deter future similar conduct;

16 d. For an order requiring Defendants to immediately cease and desist from all  
17 fraudulent, deceptive, unlawful, and illegal conduct described above;

18 e. Prejudgment interest;

19 f. Postjudgment interest;

20 g. Awarding Plaintiff's reasonable attorneys' fees;

21 h. Awarding Plaintiff the costs of these proceedings; and

22 i. Such other and further relief as this Court deems just and proper.

23 ///

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1 Dated: July 8, 2016

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3

By: /s/ Pierce Gore  
Ben F. Pierce Gore (SBN 128515)  
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CIVIL COVER SHEET

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

I. (a) PLAINTIFFS

DOLORES GOULD

(b) County of Residence of First Listed Plaintiff Contra Costa (EXCEPT IN U.S. PLAINTIFF CASES)

(c) Attorneys (Firm Name, Address, and Telephone Number) Ben F. Pierce Gore 1871 The Alameda, Suite 425 San Jose, CA 95126 (408) 429-6506

DEFENDANTS

JOHNSON & JOHNSON, and JOHNSON & JOHNSON CONSUMER COMPANIES, INC.

County of Residence of First Listed Defendant (IN U.S. PLAINTIFF CASES ONLY)

NOTE: IN LAND CONDEMNATION CASES, USE THE LOCATION OF THE TRACT OF LAND INVOLVED. Attorneys (If Known)

II. BASIS OF JURISDICTION (Place an "X" in One Box Only)

- 1 U.S. Government Plaintiff 2 U.S. Government Defendant 3 Federal Question (U.S. Government Not a Party) 4 Diversity (Indicate Citizenship of Parties in Item III)

III. CITIZENSHIP OF PRINCIPAL PARTIES (Place an "X" in One Box for Plaintiff and One Box for Defendant)

Table with columns for Plaintiff (PTF) and Defendant (DEF) citizenship options: Citizen of This State, Citizen of Another State, Citizen or Subject of a Foreign Country, Incorporated or Principal Place of Business in This State, Incorporated and Principal Place of Business in Another State, Foreign Nation.

IV. NATURE OF SUIT (Place an "X" in One Box Only)

Large table with categories: CONTRACT, REAL PROPERTY, PERSONAL INJURY, CIVIL RIGHTS, PRISONER PETITIONS, FORFEITURE/PENALTY, LABOR, IMMIGRATION, BANKRUPTCY, SOCIAL SECURITY, FEDERAL TAX SUITS, OTHER STATUTES.

V. ORIGIN (Place an "X" in One Box Only)

- 1 Original Proceeding 2 Removed from State Court 3 Remanded from Appellate Court 4 Reinstated or Reopened 5 Transferred from Another District (specify) 6 Multidistrict Litigation-Transfer 8 Multidistrict Litigation-Direct File

VI. CAUSE OF ACTION

Cite the U.S. Civil Statute under which you are filing (Do not cite jurisdictional statutes unless diversity): 28 U.S.C. 1332(d) Brief description of cause: Personal Injury due to products liability

VII. REQUESTED IN COMPLAINT:

CHECK IF THIS IS A CLASS ACTION UNDER RULE 23, Fed. R. Civ. P. DEMAND \$ CHECK YES only if demanded in complaint: JURY DEMAND: [X] Yes [ ] No

VIII. RELATED CASE(S), IF ANY (See instructions):

JUDGE DOCKET NUMBER

IX. DIVISIONAL ASSIGNMENT (Civil Local Rule 3-2)

(Place an "X" in One Box Only) [X] SAN FRANCISCO/OAKLAND [ ] SAN JOSE [ ] EUREKA-MCKINLEYVILLE

DATE: 07/08/2016

SIGNATURE OF ATTORNEY OF RECORD: /s/ Pierce Gore

## INSTRUCTIONS FOR ATTORNEYS COMPLETING CIVIL COVER SHEET FORM JS-CAND 44

**Authority For Civil Cover Sheet.** The JS-CAND 44 civil cover sheet and the information contained herein neither replaces nor supplements the filings and service of pleading or other papers as required by law, except as provided by local rules of court. This form, approved in its original form by the Judicial Conference of the United States in September 1974, is required for the Clerk of Court to initiate the civil docket sheet. Consequently, a civil cover sheet is submitted to the Clerk of Court for each civil complaint filed. The attorney filing a case should complete the form as follows:

- I. a) Plaintiffs-Defendants.** Enter names (last, first, middle initial) of plaintiff and defendant. If the plaintiff or defendant is a government agency, use only the full name or standard abbreviations. If the plaintiff or defendant is an official within a government agency, identify first the agency and then the official, giving both name and title.
- b) County of Residence.** For each civil case filed, except U.S. plaintiff cases, enter the name of the county where the first listed plaintiff resides at the time of filing. In U.S. plaintiff cases, enter the name of the county in which the first listed defendant resides at the time of filing. (NOTE: In land condemnation cases, the county of residence of the “defendant” is the location of the tract of land involved.)
- c) Attorneys.** Enter the firm name, address, telephone number, and attorney of record. If there are several attorneys, list them on an attachment, noting in this section “(see attachment).”
- II. Jurisdiction.** The basis of jurisdiction is set forth under Federal Rule of Civil Procedure 8(a), which requires that jurisdictions be shown in pleadings. Place an “X” in one of the boxes. If there is more than one basis of jurisdiction, precedence is given in the order shown below.
- (1) United States plaintiff. Jurisdiction based on 28 USC §§ 1345 and 1348. Suits by agencies and officers of the United States are included here.
  - (2) United States defendant. When the plaintiff is suing the United States, its officers or agencies, place an “X” in this box.
  - (3) Federal question. This refers to suits under 28 USC § 1331, where jurisdiction arises under the Constitution of the United States, an amendment to the Constitution, an act of Congress or a treaty of the United States. In cases where the U.S. is a party, the U.S. plaintiff or defendant code takes precedence, and box 1 or 2 should be marked.
  - (4) Diversity of citizenship. This refers to suits under 28 USC § 1332, where parties are citizens of different states. When Box 4 is checked, the citizenship of the different parties must be checked. (See Section III below; **NOTE: federal question actions take precedence over diversity cases.**)
- III. Residence (citizenship) of Principal Parties.** This section of the JS-CAND 44 is to be completed if diversity of citizenship was indicated above. Mark this section for each principal party.
- IV. Nature of Suit.** Place an “X” in the appropriate box. If the nature of suit cannot be determined, be sure the cause of action, in Section VI below, is sufficient to enable the deputy clerk or the statistical clerk(s) in the Administrative Office to determine the nature of suit. If the cause fits more than one nature of suit, select the most definitive.
- V. Origin.** Place an “X” in one of the six boxes.
- (1) Original Proceedings. Cases originating in the United States district courts.
  - (2) Removed from State Court. Proceedings initiated in state courts may be removed to the district courts under Title 28 USC § 1441. When the petition for removal is granted, check this box.
  - (3) Remanded from Appellate Court. Check this box for cases remanded to the district court for further action. Use the date of remand as the filing date.
  - (4) Reinstated or Reopened. Check this box for cases reinstated or reopened in the district court. Use the reopening date as the filing date.
  - (5) Transferred from Another District. For cases transferred under Title 28 USC § 1404(a). Do not use this for within district transfers or multidistrict litigation transfers.
  - (6) Multidistrict Litigation Transfer. Check this box when a multidistrict case is transferred into the district under authority of Title 28 USC § 1407. When this box is checked, do not check (5) above.
  - (8) Multidistrict Litigation Direct File. Check this box when a multidistrict litigation case is filed in the same district as the Master MDL docket. Please note that there is no Origin Code 7. Origin Code 7 was used for historical records and is no longer relevant due to changes in statute.
- VI. Cause of Action.** Report the civil statute directly related to the cause of action and give a brief description of the cause. **Do not cite jurisdictional statutes unless diversity.** Example: U.S. Civil Statute: 47 USC § 553. Brief Description: Unauthorized reception of cable service.
- VII. Requested in Complaint.** Class Action. Place an “X” in this box if you are filing a class action under Federal Rule of Civil Procedure 23.
- Demand. In this space enter the actual dollar amount being demanded or indicate other demand, such as a preliminary injunction.
- Jury Demand. Check the appropriate box to indicate whether or not a jury is being demanded.
- VIII. Related Cases.** This section of the JS-CAND 44 is used to identify related pending cases, if any. If there are related pending cases, insert the docket numbers and the corresponding judge names for such cases.
- IX. Divisional Assignment.** If the Nature of Suit is under Property Rights or Prisoner Petitions or the matter is a Securities Class Action, leave this section blank. For all other cases, identify the divisional venue according to Civil Local Rule 3-2: “the county in which a substantial part of the events or omissions which give rise to the claim occurred or in which a substantial part of the property that is the subject of the action is situated.”
- Date and Attorney Signature.** Date and sign the civil cover sheet.

AO 440 (Rev. 06/12) Summons in a Civil Action

UNITED STATES DISTRICT COURT

for the

Northern District of California

DOLORES GOULD,

Plaintiff(s)

v.

JOHNSON & JOHNSON, and JOHNSON & JOHNSON CONSUMER COMPANIES, INC.,

Defendant(s)

Civil Action No.

SUMMONS IN A CIVIL ACTION

To: (Defendant's name and address)

Johnson & Johnson
Johnson & Johnson Consumer Companies, Inc.

Address: One Johnson & Johnson Plaza
New Brunswick, NJ 08933

A lawsuit has been filed against you.

Within 21 days after service of this summons on you (not counting the day you received it) — or 60 days if you are the United States or a United States agency, or an officer or employee of the United States described in Fed. R. Civ. P. 12 (a)(2) or (3) — you must serve on the plaintiff an answer to the attached complaint or a motion under Rule 12 of the Federal Rules of Civil Procedure. The answer or motion must be served on the plaintiff or plaintiff's attorney, whose name and address are:

Ben F. Pierce Gore
Pratt & Associates
1871 The Alameda, Suite 425
San Jose, CA 95126
P: (408) 429-6506

If you fail to respond, judgment by default will be entered against you for the relief demanded in the complaint. You also must file your answer or motion with the court.

CLERK OF COURT

Date:

Signature of Clerk or Deputy Clerk

Civil Action No. \_\_\_\_\_

**PROOF OF SERVICE**

*(This section should not be filed with the court unless required by Fed. R. Civ. P. 4(l))*

This summons for *(name of individual and title, if any)* \_\_\_\_\_  
was received by me on *(date)* \_\_\_\_\_ .

I personally served the summons on the individual at *(place)* \_\_\_\_\_  
\_\_\_\_\_ on *(date)* \_\_\_\_\_ ; or

I left the summons at the individual's residence or usual place of abode with *(name)* \_\_\_\_\_  
\_\_\_\_\_, a person of suitable age and discretion who resides there,  
on *(date)* \_\_\_\_\_ , and mailed a copy to the individual's last known address; or

I served the summons on *(name of individual)* \_\_\_\_\_ , who is  
designated by law to accept service of process on behalf of *(name of organization)* \_\_\_\_\_  
\_\_\_\_\_ on *(date)* \_\_\_\_\_ ; or

I returned the summons unexecuted because \_\_\_\_\_ ; or

Other *(specify)*: \_\_\_\_\_ .

My fees are \$ \_\_\_\_\_ for travel and \$ \_\_\_\_\_ for services, for a total of \$ \_\_\_\_\_ .

I declare under penalty of perjury that this information is true.

Date: \_\_\_\_\_

\_\_\_\_\_  
*Server's signature*

\_\_\_\_\_  
*Printed name and title*

\_\_\_\_\_  
*Server's address*

Additional information regarding attempted service, etc: