

**BURSOR & FISHER, P.A.**

Sarah N. Westcot (State Bar No. 264916)  
701 Brickell Ave., Suite 1420  
Miami, FL 33131-2800  
Telephone: (305) 330-5512  
Facsimile: (305) 676-9006  
E-Mail: swestcot@bursor.com

**BURSOR & FISHER, P.A.**

L. Timothy Fisher (State Bar No. 191626)  
1990 North California Blvd., Suite 940  
Walnut Creek, CA 94596  
Telephone: (925) 300-4455  
Facsimile: (925) 407-2700  
E-Mail: ltfisher@bursor.com

*Counsel for Plaintiffs*

**UNITED STATES DISTRICT COURT  
NORTHERN DISTRICT OF CALIFORNIA**

JORDAN NELSON and REGINA PERALTA,  
individually and on behalf of all others similarly  
situated,

Plaintiffs,

v.

KENVUE, INC., MCNEIL CONSUMER  
HEALTHCARE, JOHNSON & JOHNSON  
CONSUMER, INC., CVS PHARMACY, INC.,  
HALEON US CAPITAL LLC, GSK PLC,  
ALBERTSONS COMPANIES, INC., TARGET  
CORPORATION, WALMART INC., and  
PERRIGO COMPANY PLC,

Defendants.

Case No.

**CLASS ACTION COMPLAINT**

JURY TRIAL DEMANDED

1 Plaintiff Jordan Nelson (“Plaintiff Nelson”) and Plaintiff Regina Peralta (“Plaintiff Peralta”)  
2 (collectively, “Plaintiffs”) bring this action on behalf of themselves and all others similarly situated  
3 against Defendants Kenvue Inc., McNeil Consumer Healthcare, Johnson & Johnson Consumer, Inc.,  
4 CVS Pharmacy, Inc., Haleon US Capital LLC, GSK Plc, Albertsons Companies, Inc., Target  
5 Corporation, Walmart Inc., and Perrigo Company Plc (collectively, “Defendants”). Plaintiffs make  
6 the following allegations pursuant to the investigation of their counsel and based upon information  
7 and belief, except as to allegations specifically pertaining to themselves and their counsel, which are  
8 based on personal knowledge.

9 **NATURE OF THE ACTION**

10 1. Nasal decongestants are over-the-counter medications that are marketed to alleviate  
11 sinus pressure and sinus congestion.

12 2. Defendants have made millions of dollars selling their nasal decongestant products.  
13 Defendants’ products include: 1) Sudafed PE Sinus Congestion; 2) Sudafed PE Head Congestion  
14 + Mucus; 3) Sudafed PE Sinus Pressure + Pain; 4) Sudafed PE Head Congestion + Pain; 5) Sudafed  
15 PE Head Congestion + Flu Severe; 6) Sudafed PE Sinus Congestion Day + Night; 7) CVS Sinus  
16 PE + Allergy; 8) CVS Sinus PE Pressure, Pain + Cold; 9) CVS Nasal Decongestant PE; 10) CVS  
17 Mucus Relief Sinus Pressure, Pain & Cough; 11) CVS Mucus Relief Sinus Severe Congestion &  
18 Pain; 12) CVS Sinus Pain & Congestion; 13) CVS Severe Sinus Pain & Congestion; 14) CVS  
19 Sinus Relief; 15) CVS Cold, Flu & Sore Throat; 16) Advil Sinus Congestion & Pain; 17) Advil  
20 Multi-Symptom Cold & Flu; 18) Advil Allergy & Congestion Relief; 19) Up & Up Sinus PE Non-  
21 Drowsy Congestion Relief; 20) Signature Care Max Strength Nasal Decongestant PE; 21) Equate  
22 Suphedrine PE; 22) Equate Sinus PE & Allergy; 23) Equate Sinus Congestion PE; 24) Tylenol  
23 Sinus + Headache Non-Drowsy; 25) Tylenol Sinus Severe; and 26) Tylenol Cold + Flu + Cough  
24 Nighttime Liquid (collectively, the “Products”).

25 3. Defendants market the Products as having the ability to provide nasal decongestant  
26 relief.

1 4. Defendants attribute the Products’ ability to provide nasal decongestion relief to the  
2 inclusion of one active ingredient: Phenylephrine (“PE”).

3 5. PE, however, is ineffective at providing nasal decongestion relief when it is taken  
4 orally.

5 6. Indeed, on September 12, 2023, an advisory panel to the U.S. Food & Drug  
6 Administration (“FDA”) unanimously agreed (16-0) that oral PE is not effective at relieving nasal  
7 congestion.

8 7. Accordingly, Defendants’ marketing and advertising concerning the Products is  
9 false, misleading, and likely to deceive the public.

10 8. Plaintiffs asserts claims on behalf of themselves and similarly situated purchasers  
11 of Defendants’ Products for violations of the California Consumers Legal Remedies Act  
12 (“CLRA”), Civil Code §§ 1750, *et seq.*, Unfair Competition Law (“UCL”), Bus. & Prof. Code §§  
13 17200, *et seq.*, False Advertising Law (“FAL”), Bus. & Prof. Code §§ 17500, *et seq.*, breach of  
14 implied warranty of merchantability, and unjust enrichment.

15 **PARTIES**

16 **Plaintiff Nelson**

17 9. Plaintiff Nelson is a resident of Pleasant Hill, California, has an intent to remain  
18 there, and is therefore a domiciliary of California.

19 10. Plaintiff Nelson purchased the 1) Sudafed PE Sinus Congestion, 2) CVS Nasal  
20 Decongestant PE, and 3) Advil Sinus Congestion & Pain products numerous times over the past  
21 three years at CVS and Walgreens stores in Walnut Creek, California. Plaintiff Nelson purchased  
22 these products to treat her nasal congestion symptoms. Before purchasing the Products, Plaintiff  
23 Nelson reviewed information about the Products, including the representation that the Products  
24 would be able to provide nasal congestion relief. When reviewing the Products label, disclosures,  
25 warranties, and marketing materials, Plaintiff Nelson understood them as representations and  
26 warranties by Defendants that the Products would be able to provide nasal decongestion relief.

1           11. Plaintiff Nelson relied on Defendants’ representations and warranties in deciding to  
2 purchase the Products over other nasal decongestant products. Accordingly, Defendants’  
3 representations and warranties were part of the basis of the bargain, in that she would not have  
4 purchased the Products on the same terms had she known Defendants’ representations were not true.

5           12. Contrary to the representations on the Products’ marketing materials, the Products  
6 were not able to provide nasal decongestion relief. Plaintiff Nelson therefore did not receive the  
7 benefit of her bargain.

8 **Plaintiff Peralta**

9           13. Plaintiff Peralta is a resident of San Jose, California, has an intent to remain there, and  
10 is therefore a domiciliary of California.

11           14. Plaintiff Peralta purchased the 1) Advil Sinus Congestion & Pain, 2) Equate  
12 Suphedrine PE, 3) Tylenol Sinus + Headache, 4) Up & Up Sinus PE Non-Drowsy Congestion Relief,  
13 and 5) Signature Care Max Strength Nasal Decongestant PE products numerous times over the past  
14 three years at local CVS, Walmart, and Safeway stores in San Jose, California. Plaintiff Peralta  
15 purchased these products to treat her nasal congestion symptoms. Before purchasing the Products,  
16 Plaintiff Peralta reviewed information about the Products, including the representations that the  
17 Products would be able to provide nasal decongestant relief. When reviewing the Products label,  
18 disclosures, warranties, and marketing materials, Plaintiff Peralta understood them as representations  
19 and warranties by Defendants that the Products would be able to provide nasal decongestion relief.

20           15. Plaintiff Peralta relied on Defendants’ representations and warranties in deciding to  
21 purchase the Products over other nasal decongestant products. Accordingly, Defendants’  
22 representations and warranties were part of the basis of the bargain, in that she would not have  
23 purchased the Products on the same terms had she known Defendants’ representations were not true.

24           16. Contrary to the representations on the Products’ marketing materials, the Products  
25 were not able to provide nasal decongestion relief. Plaintiff Peralta therefore did not receive the  
26 benefit of her bargain.

1 **Defendants**

2 17. Defendant Kenvue Inc. is an American consumer health company, and formerly the  
3 consumer healthcare division of Johnson & Johnson. Kenvue is headquartered in Skillman, New  
4 Jersey. It wholly owns Defendant McNeil Consumer Healthcare.

5 18. Defendant McNeil Consumer Healthcare is wholly owned by Defendant Kenvue,  
6 with headquarters in Fort Washington, Pennsylvania. Defendant McNeil Consumer Healthcare  
7 manufactures and markets the Products Sudafed PE Sinus Congestion, Sudafed PE Head Congestion  
8 + Mucus, Sudafed PE Sinus Pressure + Pain, Sudafed PE Head Congestion + Pain, Sudafed PE Head  
9 Congestion + Flu Severe, Sudafed PE Sinus Congestion Day + Night, Tylenol Sinus + Headache  
10 Non-Drowsy, Tylenol Sinus Severe, and Tylenol Cold + Flu + Cough Nighttime Liquid throughout  
11 the state of California and the United States.

12 19. Defendant Johnson & Johnson Consumer, Inc. is a New Jersey based medical  
13 corporation, with its headquarters in New Brunswick, New Jersey. Defendant Johnson & Johnson  
14 Consumer, Inc. manufactures, markets, and sells the Products Sudafed PE Sinus Congestion, Sudafed  
15 PE Head Congestion + Mucus, Sudafed PE Sinus Pressure + Pain, Sudafed PE Head Congestion +  
16 Pain, Sudafed PE Head Congestion + Flu Severe, Sudafed PE Sinus Congestion Day + Night,  
17 Tylenol Sinus + Headache Non-Drowsy, Tylenol Sinus Severe, and Tylenol Cold + Flu + Cough  
18 Nighttime Liquid throughout the state of California and the United States.

19 20. Defendant CVS Pharmacy, Inc. is a Rhode Island corporation with its headquarters  
20 in Woonsocket, Rhode Island. Defendant CVS Pharmacy, Inc. manufactures, markets, and sells the  
21 Products CVS Sinus PE + Allergy, CVS Sinus PE Pressure, Pain + Cold, CVS Nasal Decongestant  
22 PE, CVS Mucus Relief Sinus Pressure, Pain & Cough, CVS Mucus Relief Sinus Severe Congestion  
23 & Pain, CVS Sinus Pain & Congestion, CVS Severe Sinus Pain & Congestion, CVS Sinus Relief,  
24 and CVS Cold, Flu & Sore Throat throughout the State of California and the United States.

25 21. Defendant Haleon US Capital LLC is a Delaware limited liability company with its  
26 headquarters in Warren, New Jersey. Upon information and belief, since approximately July 2022,  
27 Defendant Haleon US Capital LLC has manufactured, marketed, and sold the Products Advil Sinus  
28

1 Congestion & Pain, Advil Multi-Symptom Cold & Flu, and Advil Allergy & Congestion Relief  
2 throughout the state of California and the United States.

3 22. Defendant GSK Plc is an English company with its U.S. headquarters in Philadelphia,  
4 Pennsylvania. Upon information and belief, prior to approximately July 2022, Defendant GSK Plc  
5 had manufactured, marketed, and sold the Products Advil Sinus Congestion & Pain, Advil Multi-  
6 Symptom Cold & Flu, and Advil Allergy & Congestion Relief throughout the state of California and  
7 the United States.

8 23. Defendant Albertsons Companies, Inc., is a Delaware company with its headquarters  
9 in Boise, Idaho. Defendant Albertsons Companies, Inc., manufactures, markets, and sells the Product  
10 Signature Care Max Strength Nasal Decongestant PE throughout the state of California and the  
11 United States.

12 24. Defendant Target Corporation is a Minnesota corporation with its principal place of  
13 business in Minneapolis, Minnesota. Defendant Target Corporation manufactures, markets, and sells  
14 the Up & Up Sinus PE Non-Drowsy Congestion Relief Product throughout the State of California  
15 and the United States.

16 25. Defendant Walmart Inc. is a Delaware corporation with its headquarters in  
17 Bentonville, Arkansas. Defendant Walmart Inc. manufactures, markets, and sells the Products  
18 Equate Suphedrine PE, Equate Sinus PE & Allergy, and Equate Sinus Congestion PE throughout the  
19 State of California and the United States.

20 26. Defendant Perrigo Company Plc is an American Irish corporation with its U.S.  
21 headquarters in Allegan, Michigan. Defendant Perrigo Company Plc manufactures, markets, and  
22 sells the Products Equate Suphedrine PE, Equate Sinus PE & Allergy, and Equate Sinus  
23 Congestion PE throughout the State of California and the United States.

24 **JURISDICTION AND VENUE**

25 27. This Court has subject matter jurisdiction pursuant to 28 U.S.C. § 1332(d)(2)(A)  
26 because this case is a class action where the aggregate claims of all members of the proposed class  
27 are in excess of \$5,000,000.00, exclusive of interest and costs, there are over 100 members of the  
28

1 putative class, and Plaintiffs, as well as most members of the proposed class, are citizens of states  
2 different than Defendants.

3 28. The Court has personal jurisdiction over Defendants because Defendants conduct  
4 substantial business within California, such that Defendants have significant, continuous, and  
5 pervasive contacts with the State of California. Moreover, Defendants have purposefully availed  
6 themselves of the laws and benefits of doing business in California, and Plaintiffs' claims arise out  
7 of the Defendants' forum-related activities.

8 29. Venue is proper in this Court pursuant to 28 U.S.C. § 1391(b) because Defendants  
9 transact significant business within this District and because Plaintiffs purchased and used the  
10 Products in this District.

### 11 **FACTUAL ALLEGATIONS**

#### 12 ***The Market For Decongestants***

13 30. The market for products that allegedly relieve nasal congestion is worth over \$2  
14 billion annually and includes over 250 products.

15 31. The two leading ingredients used to provide relief from nasal congestion are PE and  
16 pseudoephedrine. PE is the only active ingredient in the Products advertised to provide relief for  
17 nasal congestion.

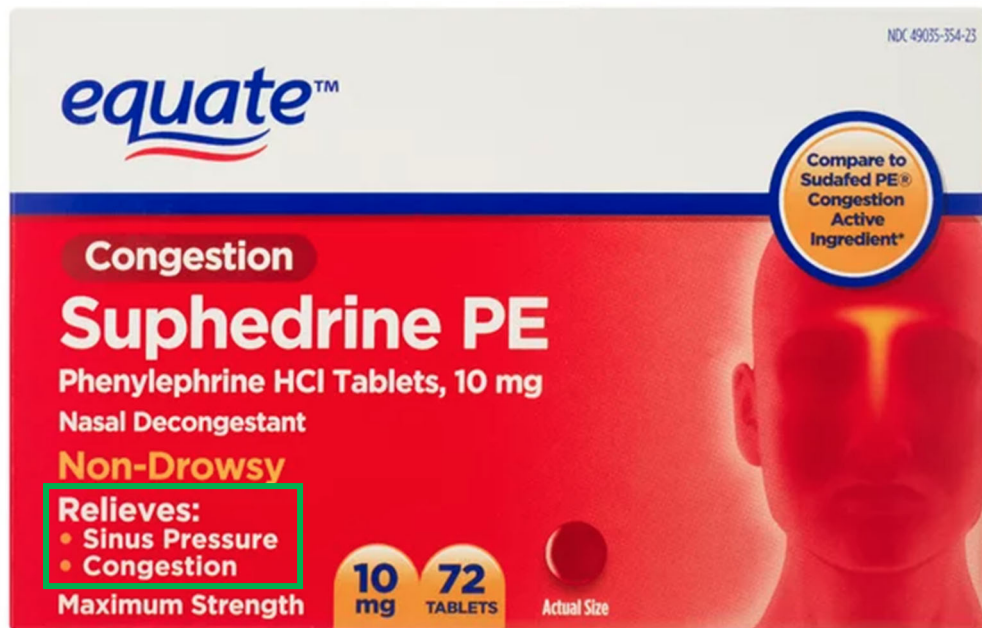
18 32. While pseudoephedrine is effective as a nasal decongestant when taken orally, PE  
19 is not. Nonetheless, PE accounts for approximately 80% of the market for over-the-counter  
20 decongestants. In the last year alone, nearly \$1.8 billion of PE-based decongestants were sold.

#### 21 ***Defendants' False Advertising***

22 33. Defendants market, sell, and distribute the Products through numerous brick-and-  
23 mortar stores as well as online. On the Products packaging, Defendants represent that the Products  
24 are able to provide nasal decongestion relief.

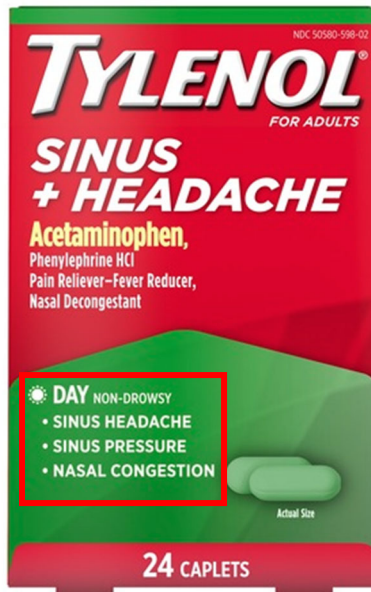
25 34. By way of example, the Products include, but are not limited to, those depicted by  
26 below:

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28

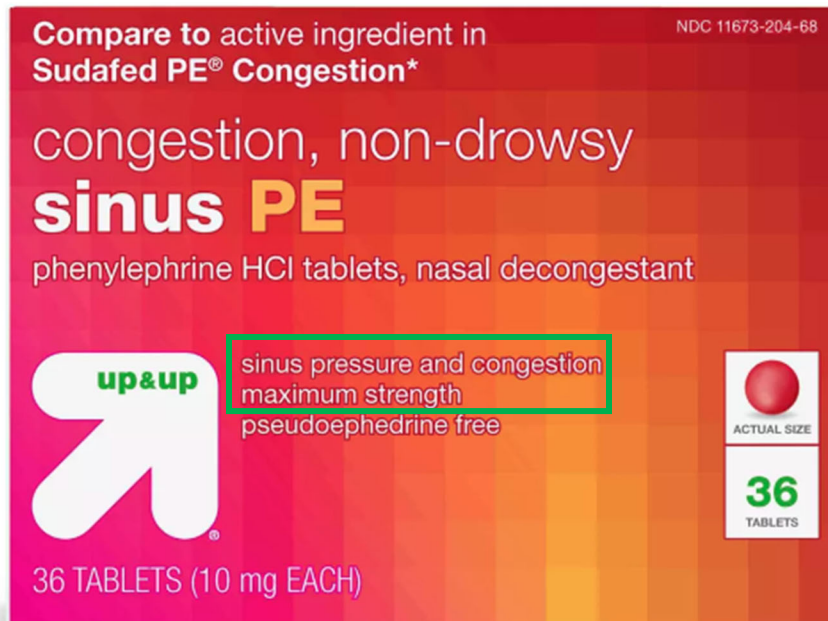




1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28



1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28





35. By representing that the Products are effective remedies for nasal congestion, Defendants induced reasonable consumers, such as Plaintiffs and the proposed class members into believing that the Products were effective at providing nasal decongestion relief. Those representations, however, are false and misleading, as set forth in greater detail below.

***The Products' Use of Phenylephrine***

36. Defendants' Products all attribute the ability to provide nasal decongestion relief to one active ingredient: PE.

37. Defendants do not attribute nasal decongestant relief to any other ingredient in the Products.

***Phenylephrine Does Not Provide Nasal Decongestant Relief When Taken Orally***

38. PE is ineffective at providing nasal decongestant relief when taken orally. All available scientific authorities support this conclusion.

39. For example, on May 1, 2006, two professors at the University of Florida, Dr. Leslie Hendeles, PharmD Professor, Pharmacy and Pediatrics, and Dr. Randy Hatton, PharmD FCCP BCPS Clinical Professor, Department of Pharmacotherapy and Translational Research College of Pharmacy published a letter in Journal of Allergy and Clinical Immunology titled: Oral

1 phenylephrine: An ineffective replacement for pseudophedrine?<sup>1</sup> The letter questioned the  
2 effectiveness of PE for nasal congestion based upon the results of multiple double blind, placebo-  
3 controlled studies, that show PE was no more effective than placebo in reducing nasal airway  
4 resistance. Moreover, the letter notes that the studies relied on by the FDA to approve PE were  
5 unpublished, manufacturer-sponsored studies conducted by commercial testing laboratories.

6 40. On February 1, 2007, three professors from the University of Florida, Dr. Leslie  
7 Hendeles, PharmD Professor, Pharmacy and Pediatrics, Dr. Randy Hatton, PharmD FCCP BCPS  
8 Clinical Professor, Department of Pharmacotherapy and Translational Research College of  
9 Pharmacy, and Almut G. Winterstein (PhD, Assistant Professor, Department of Healthcare  
10 Administration) filed a Citizens Petition with the FDA concerning PE drugs.<sup>2</sup>

11 41. As a result of the 2007 Citizens Petition, the FDA's Nonprescription Drugs  
12 Advisory Committee met on December 14, 2007 and concluded that the products could continue  
13 to be sold, but 9 of 12 of the committee members voted that "new studies on response to higher  
14 doses were required."<sup>3</sup>

15 42. Scherling-Plough Pharmaceuticals responded to the FDA's Nonprescription Drugs  
16 Advisory Committee by conducting a multicenter, phase 2, trial among 539 adults with seasonal  
17 allergic rhinitis. The results of the study revealed no significant differences between placebo and  
18 active treatment groups.<sup>4</sup>

19 43. In addition, McNeil Consumer Healthcare conducted a pharmacokinetic, safety and  
20 cardiovascular tolerability study of phenylephrine. Similarly, this study revealed no difference in  
21 safety endpoints between placebo and 10, 20, and 30 mg of phenylephrine even though systemic  
22 exposure increased disproportionately with dose. "This is noteworthy since both the relief of  
23 congestion and systemic endpoints such as change in blood pressure and pulse are mediated by

24 \_\_\_\_\_  
25 <sup>1</sup> [https://www.jacionline.org/article/S0091-6749\(06\)00633-6/fulltext](https://www.jacionline.org/article/S0091-6749(06)00633-6/fulltext) (last accessed Sept. 18, 2023).

26 <sup>2</sup> <https://www.regulations.gov/docket/FDA-2007-P-0108/document> (last accessed Sept. 18, 2023).

27 <sup>3</sup> [https://www.jaci-inpractice.org/article/S2213-2198\(15\)00318-9/fulltext](https://www.jaci-inpractice.org/article/S2213-2198(15)00318-9/fulltext) (last accessed Sept. 18,  
28 2023).

<sup>4</sup> [chrome-extension://efaidnbmninnibpcapjpcgkclefindmkaj/https://truthinadvertising.org/wp-  
content/uploads/2023/02/Hatton-Hendeles-2015-Citizens-Petition-re-oral-phenylephrine.pdf](chrome-extension://efaidnbmninnibpcapjpcgkclefindmkaj/https://truthinadvertising.org/wp-content/uploads/2023/02/Hatton-Hendeles-2015-Citizens-Petition-re-oral-phenylephrine.pdf) (last  
accessed Sept. 15, 2023).

1 alpha adrenergic stimulation. The absence of a significant effect on the latter at the higher doses  
2 suggest that the concentrations reached are not sufficient to stimulate alpha adrenergic receptors.”<sup>5</sup>

3 44. On November 4, 2015, another Citizens Petition was filed by two professors at the  
4 University of Florida, Dr. Leslie Hendeles, PharmD Professor, Pharmacy and Pediatrics, and Dr.  
5 Randy Hatton, PharmD FCCP BCPS Clinical Professor, Department of Pharmacotherapy and  
6 Translational Research College of Pharmacy. The petition asked the FDA “to remove oral  
7 phenylephrine from the Final Monograph for OTC nasal decongestant products.”<sup>6</sup> Specifically,  
8 the petition asked the FDA to remove Phenylephrine and to remove phenylephrine bitartrate, “both  
9 individually and in combination drug products in an effervescent dosage form.”<sup>7</sup>

10 45. According to the 2015 Citizens Petition, “[t]wo additional studies published in 2009  
11 provide further evidence of the absence of a decongestant effect from the FDA-approved  
12 nonprescription does of 10mg,” and “PE was not significantly different from placebo in the mean  
13 change in subjective nasal congestion scores whereas pseudoephedrine, a positive control in the  
14 study, decreased congestion significantly greater than placebo and PE.”<sup>8</sup>

15 ***The FDA Advisory Panel’s Recent Vote on PE***

16 46. Recently, “[t]he FDA held a Non-prescription Drug Advisory Committee meeting  
17 ... to discuss the effectiveness of oral phenylephrine as an active ingredient in over-the-counter  
18 (OTC) cough and cold products that are indicated for the temporary relief of congestion, both as a  
19 single ingredient product and in combination with other ingredients.”<sup>9</sup>

20 47. In doing so, the Panel referenced numerous studies demonstrating that PE is not  
21 effective for treating nasal congestion when taken orally.

22 48. As a result, the Panel concluded that “the current scientific data do[es] not support  
23 that the recommended dosage of orally administered phenylephrine is effective as a nasal  
24

---

25 <sup>5</sup> *Id.*

26 <sup>6</sup> <https://truthinadvertising.org/wp-content/uploads/2023/02/Hatton-Hendeles-2015-Citizens-Petition-re-oral-phenylephrine.pdf> (last accessed Sept. 18, 2023).

27 <sup>7</sup> *Id.*

28 <sup>8</sup> *Id.*

<sup>9</sup> <https://www.fda.gov/drugs/drug-safety-and-availability/fda-clarifies-results-recent-advisory-committee-meeting-oral-phenylephrine> (last accessed Sept. 15, 2023).

1 decongestant.”<sup>10</sup>

2 49. In fact, the Panel members voted unanimously (16-0) that PE drugs were ineffective  
3 when taken orally.

4 ***Misbranded Drugs Are Illegal to Sell***

5 50. As OTC drug products regulated by the FDA, the Products must be both safe *and*  
6 effective and are subject to federal current Good Manufacturing Practices (“cGMP”) regulations and  
7 the FDCA’s state law analogues. These cGMP regulations require OTC medications like the  
8 Products to meet safety, quality, purity, identity, and strength standards. *See* 21 U.S.C.  
9 § 351(a)(2)(B).

10 51. The cGMPs establish “minimum current good manufacturing practice for methods to  
11 be used in, and the facilities or controls to be used for, the manufacture, processing, packing, or  
12 holding of a drug to assure that such drug meets the requirements of the act as to safety, and has the  
13 identity and strength and meets the quality and purity characteristics that it purports or is represented  
14 to possess.” 21 C.F.R. § 210.1(a). In other words, manufacturers, like Defendants, at all phases of  
15 the design, manufacture, and distribution chain are bound by these requirements.

16 52. The cGMPs set forth minimum standards regarding: organization and personnel  
17 (Subpart B); buildings and facilities (Subpart C); equipment (Subpart D); control of components and  
18 drug product containers and closures (Subpart E); production and process controls (Subpart F);  
19 packaging and label controls (Subpart G); holding and distribution (Subpart H); laboratory controls  
20 (Subpart I); records and reports (Subpart J); and returned and salvaged drug products (Subpart K).  
21 The FDA has worldwide jurisdiction to enforce these regulations if the facility is making drugs  
22 intended to be distributed in the United States.

23 53. Any drug product not manufactured in accordance with cGMPs is deemed  
24 “adulterated” or “misbranded” and may not be distributed or sold in the United States. *See* 21 U.S.C.  
25 §§ 331(a), 351(a)(2)(B). States have enacted laws adopting or mirroring these federal standards.

26  
27 <sup>10</sup> [https://www.fda.gov/drugs/drug-safety-and-availability/fda-clarifies-results-recent-advisory-](https://www.fda.gov/drugs/drug-safety-and-availability/fda-clarifies-results-recent-advisory-committee-meeting-oral-phenylephrine)  
28 [committee-meeting-oral-phenylephrine](https://www.fda.gov/drugs/drug-safety-and-availability/fda-clarifies-results-recent-advisory-committee-meeting-oral-phenylephrine) (last accessed Sept. 15, 2023).

1           54.     FDA regulations require a drug product manufacturer to have “written procedures for  
2 production and process control designed to assure that the drug products have the identity, strength,  
3 quality, and purity they purport or are represented to possess.” 21 C.F.R. § 211.100.

4           55.     A drug product manufacturer’s “[l]aboratory controls shall include the establishment  
5 of scientifically sound and appropriate specifications, standards, sampling plans, and test procedures  
6 designed to assure that components, drug product containers, closures, in-process materials, labeling,  
7 and drug products conform to appropriate standards of identity, strength, quality, and purity.” 21  
8 C.F.R. § 211.160.

9           56.     “Laboratory records shall include complete data derived from all tests necessary to  
10 assure compliance with established specifications and standards, including examinations and assays”  
11 and a “statement of the results of tests and how the results compare with established standards of  
12 identity, strength, quality, and purity for the component, drug product container, closure, in-process  
13 material, or drug product tested.” 21 C.F.R. § 211.194(a)(6).

14           57.     Defendants could have avoided any potential for misrepresenting the quality  
15 characteristics that it represented the Products possessed by testing the effectiveness of PE in the  
16 Products for the purported claims on the Products’ labeling.

17           58.     The ineffectiveness of PE in the Products renders the Products both adulterated and  
18 misbranded under the FDCA. The Products are adulterated because they are “drug[s] and the  
19 methods used in, or the facilities or controls used for, its manufacture, processing, packing, or holding  
20 do not conform to or are not operated or administered in conformity with current good manufacturing  
21 practice to assure that such drug meets the requirements of this chapter as to safety and has the  
22 identity and strength, and meets the quality and purity characteristics, which it purports or is  
23 represented to possess.” 21 U.S.C. § 351(a)(1).

24           59.     The Products are misbranded because their labeling is “false” and “misleading”  
25 because it does not alleviate nasal congestion and/or sinus relief. 21 U.S.C. § 352(a)(1).

1           60. A product that is “adulterated” or “misbranded” cannot legally be manufactured,  
2 advertised, distributed, or sold. 21 U.S.C. § 331(a). Adulterated and misbranded products thus have  
3 no economic value and are legally worthless.

4           61. As alleged herein, Defendants have violated the FDCA, California’s Consumers  
5 Legal Remedies Act (“CLRA”), California’s Unfair Competition Law (“UCL”), California’s False  
6 Advertising Law (“FAL”), and other consumer protection statutes. Defendants engaged in  
7 fraudulent, unfair, deceptive, misleading, and/or unlawful conduct stemming from its  
8 misrepresentations and omissions surrounding the quality and purity characteristics affecting the  
9 Products.

10           62. If Defendants had disclosed to Plaintiffs and putative Class Members that the  
11 Products do not have the quality characteristics that it purports or is represented to possess, Plaintiffs  
12 and putative Class Members would not have purchased the Products  
13 or they would have paid less for the Products.

14           63. As a seller of an OTC drug product, Defendants had and has a duty to ensure that its  
15 Products have the identity and strength and meets the quality characteristics that it purports or is  
16 represented to possess, including through regular testing, especially before the Products are injected  
17 into the stream of commerce for consumers to use on their bodies. But based on the FDA Panel’s  
18 conclusions set forth above, Defendants made no reasonable effort to test its Products for the nasal  
19 decongestant claims it made. Nor did it disclose to Plaintiffs in any advertising or marketing that the  
20 Products did not conform to the nasal decongestant claims it purported or represented to possess. To  
21 the contrary, Defendants represented and warranted, expressly and impliedly, that the Products were  
22 of merchantable quality, complied with federal and state law, and did have the identity and strength  
23 and meet the quality characteristics that it purports or is represented to possess.

24 ***Injuries to Plaintiffs and Class Members***

25           64. When Plaintiffs purchased Defendants’ Products, Plaintiffs did not know, and had no  
26 reason to know, that Defendants’ Products did not have the identity and strength and meet the quality  
27 characteristics that it purported to possess (*i.e.*, the ability to alleviate nasal congestion). Not only  
28



1 would Plaintiffs not have purchased Defendants' Products had Plaintiffs known the Products did not  
2 have the ability to alleviate nasal congestion, but Plaintiffs would also not have been capable of  
3 purchasing them if Defendants had done as the law required and tested the Products for its ability to  
4 alleviate nasal congestion.

5 65. Consumers lack the ability to test or independently ascertain or verify whether a  
6 product has the identity and strength and meets the quality characteristics that it purports to possess,  
7 especially at the point of sale, and therefore must rely on Defendants to truthfully and honestly report  
8 what the Products can do on the Products' packaging or labels.

9 66. Further, given Defendants' position in the health and medication market as an  
10 industry leader, Plaintiffs and reasonable consumers trusted and relied on Defendants'  
11 representations and omissions regarding the ability to alleviate nasal congestion in the Products.

12 67. Yet, when consumers look at the Products' packaging, the Products are represented  
13 as having the ability to alleviate nasal congestion. This leads reasonable consumers to believe the  
14 Products have the ability to alleviate nasal congestion.

15 68. No reasonable consumer would have paid any amount for products that do not have  
16 the ability to alleviate nasal congestion, when the Products are marketed to consumers as having the  
17 ability to alleviate nasal congestion.

18 69. Thus, if Plaintiffs and Class members had been informed that Defendants' Products  
19 do not have the ability to alleviate nasal congestion, they would not have purchased or used the  
20 Products, or would have paid significantly less for the Products, making such omitted facts material  
21 to them.

22 70. Defendant's false, misleading, omissions, and deceptive misrepresentations regarding  
23 the Products' ability to alleviate nasal congestion are likely to continue to deceive and mislead  
24 reasonable consumers and the public, as it has already deceived and misled Plaintiffs and the Class  
25 Members.

26 71. Plaintiffs and Class members bargained for a Product that has the ability to alleviate  
27 nasal congestion. Plaintiffs and Class members were injured by the full purchase price of the  
28

1 Products because the Products are worthless, as they do not have the ability to alleviate nasal  
2 congestion, and Defendants failed to warn consumers of this fact. Such illegally sold products are  
3 worthless and have no value.

4 72. As alleged above, Plaintiffs and Class members' Products do not have the ability to  
5 alleviate nasal congestion, despite the Products' representations to the contrary.

6 73. Plaintiffs and Class members are further entitled to statutory and punitive damages,  
7 attorneys' fees and costs, and any further relief this Court deems just and proper.

8 **CLASS ALLEGATIONS**

9 74. Plaintiffs, individually and on behalf of all others, brings this class action pursuant to  
10 Fed. R. Civ. P. 23.

11 75. Plaintiffs seek to represent a class defined as:

12 All persons who purchased one or more of Defendants' Products in the  
13 United States for personal or household use within any applicable  
14 limitations period ("Nationwide Class").

15 76. Plaintiffs also seek to represent a subclass defined as:

16 All persons who purchased one or more of Defendants' Products in  
17 California for personal or household use within any applicable  
18 limitations period ("California Subclass").

19 77. Excluded from the Class and Subclass are: (1) any Judge or Magistrate presiding over  
20 this action and any members of their families; (2) Defendants, Defendants' subsidiaries, parents,  
21 successors, predecessors, and any entities in which Defendants or its parents and any entities in which  
22 Defendants have a controlling interest and its current or former employees, officers, and directors;  
23 and (3) individuals who allege personal bodily injury resulting from the use of the Products.

24 78. Plaintiffs reserve the right to modify, change, or expand the definitions of the Class  
25 and/or Subclass based upon discovery and further investigation.

26 79. *Numerosity*: The Class is so numerous that joinder of all members is impracticable.  
27 The Class likely contains thousands of members based on publicly available data. The Class is  
28 ascertainable by records in Defendants' possession.

1           80.    *Commonality*: Questions of law or fact common to the Class include, without  
2 limitation:

- 3                   • Whether the Products have the ability to alleviate nasal congestion;
- 4                   • Whether a reasonable consumer would consider the Products inability to alleviate  
5 nasal congestion to be material;
- 6                   • Whether Defendants knew or should have known that the Products do not have  
7 the ability to alleviate nasal congestion;
- 8                   • Whether Defendants misrepresented whether the Products have the ability to  
9 alleviate nasal congestion;
- 10                  • Whether Defendants failed to disclose that the Products do not have the ability to  
11 alleviate nasal congestion;
- 12                  • Whether Defendants concealed that the Products do not have the ability to  
13 alleviate nasal congestion;
- 14                  • Whether Defendants engaged in unfair or deceptive trade practices;
- 15                  • Whether Defendants violated the state consumer protection statutes alleged  
16 herein;
- 17                  • Whether Defendants were unjustly enriched; and
- 18                  • Whether Plaintiffs and Class members are entitled to damages.

19           81.    *Typicality*: Plaintiffs' claims are typical of the claims of Class members. Plaintiffs  
20 and Class members were injured and suffered damages in substantially the same manner, have the  
21 same claims against Defendants relating to the same course of conduct, and are entitled to relief  
22 under the same legal theories.

23           82.    *Adequacy*: Plaintiffs will fairly and adequately protect the interests of the Class and  
24 have no interests antagonistic to those of the Class. Plaintiffs have retained counsel experienced in  
25 the prosecution of complex class actions, including actions with issues, claims, and defenses similar  
26 to the present case. Counsel intends to vigorously prosecute this action.

1 83. *Predominance and superiority*: Questions of law or fact common to Class members  
 2 predominate over any questions affecting individual members. A class action is superior to other  
 3 available methods for the fair and efficient adjudication of this case because individual joinder of all  
 4 Class members is impracticable and the amount at issue for each Class member would not justify the  
 5 cost of litigating individual claims. Should individual Class members be required to bring separate  
 6 actions, this Court would be confronted with a multiplicity of lawsuits burdening the court system  
 7 while also creating the risk of inconsistent rulings and contradictory judgments. In contrast to  
 8 proceeding on a case-by-case basis, in which inconsistent results will magnify the delay and expense  
 9 to all parties and the court system, this class action presents far fewer management difficulties while  
 10 providing unitary adjudication, economies of scale and comprehensive supervision by a single court.  
 11 Plaintiff is unaware of any difficulties that are likely to be encountered in the management of this  
 12 action that would preclude its maintenance as a class action.

13 84. Accordingly, this class action may be maintained pursuant to Fed. R. Civ. P. 23(b)(3).

14 **COUNT I**  
 15 **VIOLATIONS OF THE CALIFORNIA UNFAIR COMPETITION LAW (“UCL”)**  
**Cal. Bus. & Prof. Code § 17200, et seq.**  
 16 **(On behalf of Plaintiffs and the California Subclass)**

17 85. Plaintiffs repeat and reallege each and every allegation contained in the foregoing  
 18 paragraphs as if fully set forth herein.

19 86. Plaintiffs bring this Count on behalf of themselves and the California Subclass against  
 20 Defendants.

21 87. The UCL prohibits any “unlawful, unfair or fraudulent business act or practice and  
 22 unfair, deceptive, untrue or misleading advertising...” Cal. Bus. & Prof. Code § 17200.

23 ***Fraudulent Acts and Practices***

24 88. Any business act or practice that is likely to deceive members of the public constitutes  
 25 a fraudulent business act or practice under the UCL. Similarly, any advertising that is deceptive,  
 26 untrue or misleading constitutes a fraudulent business act or practice under the UCL.

27  
 28

1 89. Defendants have engaged in conduct that is likely to deceive members of the public.  
2 This conduct includes representing on its Products' labels that its Products have the ability to  
3 alleviate nasal congestion.

4 90. As alleged above, Defendants have engaged in deceptive, untrue, and misleading  
5 advertising by making representations regarding the quality of the Products and material omissions  
6 regarding the Products' ability to alleviate nasal congestion.

7 91. Plaintiffs and the putative Class members were exposed to one or more of these  
8 representations and/or omissions during the class period and relied on one or more of these  
9 representations and/or omissions in deciding to purchase Defendants' Products. Indeed, although  
10 the Products were found to not have the ability to alleviate nasal congestion, Defendants make  
11 representations on the Products' packaging and labels to the contrary. Again, such  
12 misrepresentations and omissions mislead consumers regarding the quality of the Products.

13 92. By committing the acts alleged above, Defendants have engaged in fraudulent  
14 business acts and practices, which constitute unfair competition within the meaning of Business &  
15 Professions Code §17200.

16 ***Unlawful Acts and Practices***

17 93. The violation of any law constitutes an unlawful business practice under Business &  
18 Professions Code §17200.

19 94. Defendants' conduct also violates Cal. Health & Safety Code § 111730, which  
20 prohibits the sale of any misbranded product. By selling Products that do not accurately reflect the  
21 quality of the Products, the labeling is "false and misleading in any particular" in violation of Health  
22 & Safety Code § 111730.

23 95. By violating Cal. Health and Safety Code § 111730, Defendants have engaged in  
24 unlawful business acts and practices which constitute unfair competition within the meaning of Cal.  
25 Bus. & Prof. Code § 17200.

*Unfair Acts and Practices*

1  
2 96. Any business practice that offends an established public policy or is immoral,  
3 unethical, oppressive, unscrupulous, or substantially injurious to consumers constitutes an “unfair”  
4 practice under the UCL.

5 97. Defendants have engaged in unfair business practices. This conduct includes  
6 representing that the Products have the ability to alleviate nasal congestion.

7 98. Defendants have engaged in conduct that violates the legislatively declared policies  
8 of the FTC Act against committing unfair methods of competition and unfair or deceptive acts or  
9 practices in or affecting commerce. Defendants gained an unfair advantage over its competitors,  
10 whose advertising for products must comply with the FTC Act.

11 99. Defendants’ conduct, including misrepresenting the qualities of the Products, is  
12 substantially injurious to consumers. Plaintiffs and the Class would not have paid for nasal  
13 decongestant products that do not have the ability to alleviate nasal congestion but for Defendants’  
14 false labeling, advertising, and promotion. Thus, Plaintiffs and the putative Class have “lost money  
15 or property” as required for UCL standing, and such an injury is not outweighed by any  
16 countervailing benefits to consumers or competition.

17 100. Indeed, no benefit to consumers or competition results from Defendants’ conduct.  
18 Since consumers reasonably rely on Defendants’ representation of the qualities described in the  
19 Products’ labels and injury resulted from ordinary use of the Products, consumers could not have  
20 reasonably avoided such injury.

21 101. By committing the acts described above, Defendants have engaged in unfair business  
22 acts and practices which constitute unfair competition within the meaning of the UCL.

23 102. As a result of the conduct described above, Defendants have been unjustly enriched  
24 at the expense of the Plaintiffs and the putative Class.

25 103. An action for restitution is specifically authorized under Cal. Bus. & Prof. Code  
26 17203.

1           104. Wherefore, Plaintiffs pray for judgment against Defendants, as set forth hereafter.  
2 Defendants' conduct with respect to the labeling, advertising, marketing, and sale of the Products is  
3 unfair because Defendants' conduct was immoral, unethical, unscrupulous, or substantially injurious  
4 to consumers and the utility of its conduct, if any, does not outweigh the gravity of the harm to its  
5 victims.

6           105. On behalf of Plaintiffs and the putative Class, Plaintiffs seek an order for the  
7 restitution of all monies spent on the Products, which were acquired through acts of fraudulent,  
8 unfair, or unlawful competition. In addition, because the Products admittedly do not have the ability  
9 to alleviate nasal congestion, the measure of restitution should be rescission and full refund insofar  
10 as the Products are worthless. But for Defendants' misrepresentations and omissions, Plaintiffs  
11 would have paid nothing for Products that do not have the ability to alleviate nasal congestion.  
12 Indeed, there is no discernible "market" for an OTC nasal decongestant that does not have the ability  
13 to alleviate nasal congestion. As a result, the Products are rendered valueless.

14           106. Plaintiffs and California Subclass Members have no adequate remedy at law for this  
15 claim. Plaintiffs plead their claim for equitable relief in the alternative, which inherently would  
16 necessitate a finding of no adequate remedy at law.

17           107. Alternatively, legal remedies available to Plaintiffs are inadequate because they are  
18 not "equally prompt and certain and in other ways efficient" as equitable relief. *American Life Ins.*  
19 *Co. v. Stewart*, 300 U.S. 203, 214 (1937); *see also United States v. Bluit*, 815 F. Supp. 1314, 1317  
20 (N.D. Cal. Oct. 6, 1992) ("The mere existence' of a possible legal remedy is not sufficient to  
21 warrant denial of equitable relief."); *Quist v. Empire Water Co.*, 2014 Cal. 646, 643 (1928) ("The  
22 mere fact that there may be a remedy at law does not oust the jurisdiction of a court of equity. To  
23 have this effect, the remedy must also be speedy, adequate, and efficacious to the end in view ... It  
24 must reach the whole mischief and secure the whole right of the party in a perfect manner at the  
25 present time and not in the future.").

**COUNT II**  
**VIOLATIONS OF THE CALIFORNIA FALSE ADVERTISING LAW**  
**Cal. Bus. & Prof. Code § 17500, et seq.**  
**(On behalf of Plaintiffs and the California Subclass)**

1  
2  
3 108. Plaintiffs repeat and reallege each and every allegation contained in the foregoing  
4 paragraphs as if fully set forth herein.

5 109. Plaintiffs bring this Count on behalf of themselves and the California Subclass against  
6 Defendants.

7 110. California’s False Advertising Law prohibits any statement in connection with the  
8 sale of goods “which is untrue or misleading.” Cal. Bus. & Prof. Code §17500.

9 111. As set forth herein, Defendants’ marketing claims that its Products are able to provide  
10 nasal decongestion relief are untrue and misleading. To the contrary, the Products do not have the  
11 ability to alleviate nasal congestion.

12 112. Defendants knew, or reasonably should have known, that their claims regarding the  
13 quality of its Products and/or omissions regarding the Products inability to alleviate nasal congestion  
14 were untrue or misleading.

15 113. Plaintiffs and members of the California Subclass are entitled to monetary relief, and  
16 restitution in the amount they spent on the Products.

17 114. Plaintiffs and California Subclass Members have no adequate remedy at law for this  
18 claim. Plaintiffs plead their claim for equitable relief in the alternative, which inherently would  
19 necessitate a finding of no adequate remedy at law.

20 115. Alternatively, legal remedies available to Plaintiffs are inadequate because they are  
21 not “equally prompt and certain and in other ways efficient” as equitable relief. *American Life Ins.*  
22 *Co. v. Stewart*, 300 U.S. 203, 214 (1937); *see also United States v. Bluit*, 815 F. Supp. 1314, 1317  
23 (N.D. Cal. Oct. 6, 1992) (“The mere existence’ of a possible legal remedy is not sufficient to  
24 warrant denial of equitable relief.”); *Quist v. Empire Water Co.*, 2014 Cal. 646, 643 (1928) (“The  
25 mere fact that there may be a remedy at law does not oust the jurisdiction of a court of equity. To  
26 have this effect, the remedy must also be speedy, adequate, and efficacious to the end in view ... It  
27  
28



1 must reach the whole mischief and secure the whole right of the party in a perfect manner at the  
2 present time and not in the future.”).

3 **COUNT III**  
4 **VIOLATIONS OF THE CALIFORNIA CONSUMERS LEGAL REMEDIES ACT**  
5 **Cal. Bus. & Prof. Code § 1750, et seq.**  
6 **(On behalf of Plaintiffs and the California Subclass)**

7 116. Plaintiffs repeat and reallege each and every allegation contained in the foregoing  
8 paragraphs as if fully set forth herein.

9 117. Plaintiffs bring this Count on behalf of themselves and the California Subclass against  
10 Defendants.

11 118. Defendants have employed or committed methods, acts, or practices declared  
12 unlawful by Cal. Civ. Code §1770 in connection with the Products.

13 119. In particular, by failing to inform consumers that the Products do not have the ability  
14 to alleviate nasal congestion, Defendants have violated the following provisions under California  
15 Civil Code § 1770(a):

16 (5) by representing that the Products have characteristics, uses and/or  
17 benefits which they do not;

18 (7) by representing that the Products were of a particular standard,  
19 quality, or grade which they are not; and

20 (9) by advertising the Products with intent not to sell them as  
21 advertised.

22 120. Plaintiffs and California Subclass Members have no adequate remedy at law for this  
23 claim. Plaintiffs plead their claim for equitable relief in the alternative, which inherently would  
24 necessitate a finding of no adequate remedy at law.

25 121. Alternatively, legal remedies available to Plaintiffs are inadequate because they are  
26 not “equally prompt and certain and in other ways efficient” as equitable relief. *American Life Ins.*  
27 *Co. v. Stewart*, 300 U.S. 203, 214 (1937); *see also United States v. Bluit*, 815 F. Supp. 1314, 1317  
28 (N.D. Cal. Oct. 6, 1992) (“The mere existence’ of a possible legal remedy is not sufficient to  
warrant denial of equitable relief.”); *Quist v. Empire Water Co.*, 2014 Cal. 646, 643 (1928) (“The

1 mere fact that there may be a remedy at law does not oust the jurisdiction of a court of equity. To  
2 have this effect, the remedy must also be speedy, adequate, and efficacious to the end in view ... It  
3 must reach the whole mischief and secure the whole right of the party in a perfect manner at the  
4 present time and not in the future.”).

5 122. Wherefore, Plaintiffs, on behalf of themselves and all other members of the Class  
6 seek to enjoin the unlawful acts and practices described herein. Plaintiffs reserve the right to  
7 request amendment of this complaint to include a request for damages under the CLRA after  
8 complying with Civil Code 1782(a).

9 **COUNT IV**  
10 **BREACH OF IMPLIED WARRANTY OF MERCHANTABILITY**  
11 **(On behalf of Plaintiffs and the Nationwide Class)**

12 123. Plaintiffs repeat and reallege each and every allegation contained in the foregoing  
13 paragraphs as if fully set forth herein.

14 124. Plaintiffs bring this claim on behalf of themselves and the Nationwide Class against  
15 Defendants.

16 125. At all times relevant all fifty States and the District of Columbia and Puerto Rico have  
17 codified and adopted the provisions of the Uniform Commercial Code governing the implied  
18 warranty of merchantability and fitness for ordinary purpose.

19 126. Defendants were at all times a “merchant” within the meaning of Article 2 of the  
20 U.C.C., as codified under applicable law.

21 127. The Products are and were goods within the meaning of Article 2 of the U.C.C., as  
22 codified under applicable law.

23 128. Defendants were obligated to provide Plaintiffs and the other Class Members  
24 Products that were of merchantable quality, were reasonably fit for the purpose for which they were  
25 sold, and confirmed to the standards of the trade.

26 129. Defendants impliedly warranted that those drugs were of merchantable quality and fit  
27 for that purpose.  
28

1 130. Defendants breached their implied warranties, because the Products were not of  
2 merchantable quality or fit for their ordinary purpose.

3 131. Defendants' breaches of implied warranties were a direct and proximate cause of  
4 Plaintiffs' and the other Class members' damages.

5 **COUNT V**  
6 **UNJUST ENRICHMENT**  
7 **(On behalf of Plaintiffs and the Nationwide Class)**

8 132. Plaintiffs repeat and reallege each and every allegation contained in the foregoing  
9 paragraphs as if fully set forth herein.

10 133. Plaintiffs bring this Count on behalf of themselves and the Nationwide Class against  
11 Defendants.

12 134. This claim is brought under the laws of the State of California.

13 135. Defendants' conduct violated, *inter alia*, state and federal law by manufacturing,  
14 advertising, marketing, and selling the Products while misrepresenting and omitting material facts.

15 136. Defendants' unlawful conduct allowed Defendants to knowingly realize substantial  
16 revenues from selling the Products at the expense of, and to the detriment or impoverishment of,  
17 Plaintiffs and Class members and to Defendants' benefit and enrichment. Defendants have thereby  
18 violated fundamental principles of justice, equity, and good conscience.

19 137. Plaintiffs and Class members conferred significant financial benefits and paid  
20 substantial compensation to Defendants for the Products, which were not as Defendants represented  
21 them to be.

22 138. Defendants knowingly received and enjoyed the benefits conferred on it by Plaintiffs  
23 and Class members.

24 139. It is inequitable for Defendants to retain the benefits conferred by Plaintiffs and Class  
25 members' overpayments.

26 140. Plaintiffs and Class members seek establishment of a constructive trust from which  
27 Plaintiffs and Class members may seek restitution.  
28

**PRAYER FOR RELIEF**

WHEREFORE, Plaintiffs, individually and on behalf of all others similarly situated, pray for relief and judgment against Defendants as follows:

- Certifying the Class pursuant to Rule 23 of the Federal Rules of Civil Procedure, appointing Plaintiffs as representatives of the Class and Subclasses, and designating Plaintiffs' counsel as Class Counsel;
- Awarding Plaintiffs and Class members compensatory damages, in an amount to be determined at trial;
- Awarding Plaintiffs and Class members appropriate relief, including but not limited to actual damages;
- For restitution and disgorgement of profits;
- Awarding Plaintiffs and Class members reasonable attorneys' fees and costs as allowable by law;
- Awarding pre-judgment and post-judgment interest;
- For punitive damages; and
- Granting any other relief as this Court may deem just and proper.

**DEMAND FOR JURY TRIAL**

Plaintiffs hereby demand a jury trial on all issues so triable as of right.

Dated: September 22, 2023

Respectfully submitted,

**BURSOR & FISHER, P.A.**

By:           /s/ Sarah N. Westcot            
Sarah N. Westcot

Sarah N. Westcot (State Bar No. 264916)  
701 Brickell Ave., Suite 1420  
Miami, FL 33131-2800  
Telephone: (305) 330-5512  
Facsimile: (305) 676-9006  
E-Mail: swestcot@bursor.com

**BURSOR & FISHER, P.A.**  
L. Timothy Fisher (State Bar No. 191626)

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28

1990 North California Blvd., Suite 940  
Walnut Creek, CA 94596  
Telephone: (925) 300-4455  
Facsimile: (925) 407-2700  
E-Mail: ltfisher@bursor.com

*Counsel for Plaintiffs*

1                    **CLRA Venue Declaration Pursuant to California Civil Code Section 1780(d)**

2                    I, Sarah N. Westcot, declare as follows:

3                    1.            I am an attorney at law licensed to practice in the State of California and a member  
4 of the bar of this Court. I am a partner at Bursor & Fisher, P.A., counsel of record for Plaintiffs.  
5 Plaintiff Nelson resides in Pleasant Hill, California. Plaintiff Peralta resides in San Jose, California.  
6 I have personal knowledge of the facts set forth in this declaration and, if called as a witness, I could  
7 and would competently testify thereto under oath.

8                    2.            The Complaint filed in this action is filed in the proper place for trial under Civil Code  
9 Section 1780(d) in that a substantial portion of the events alleged in the Complaint occurred in the  
10 Northern District of California, as Plaintiffs purchased their Products within this District.  
11 Additionally, Defendants advertised, marketed, manufactured, distributed, and/or sold the Products  
12 at issue to Class Members from this District.

13                    I declare under the penalty of perjury under the laws of the State of California and the United  
14 States that the foregoing is true and correct and that this declaration was executed at Miami, Florida  
15 this 22nd day of September, 2023.

16  
17                    /s/ Sarah N. Westcot  
18                    Sarah N. Westcot

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

JORDAN NELSON and REGINA PERALTA, individually and on behalf of all others similarly situated,

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

(c) Attorneys (Firm Name, Address, and Telephone Number)

Sarah N. Westcot, Bursor & Fisher, P.A., 701 Brickell Ave., Suite 1420, Miami, FL 33131-2800 Tel.: (305) 330-5512

DEFENDANTS

KENVUE, INC., MCNEIL CONSUMER HEALTHCARE, JOHNSON & JOHNSON CONSUMER, INC., CVS PHARMACY, INC., HALEON US CAPITAL LLC, GSK PLC, ALBERTSONS COMPANIES, INC., TARGET CORPORATION, WALMART INC., and PERRIGO COMPANY PLC, 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 3 Federal Question (U.S. Government Not a Party) 2 U.S. Government Defendant 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 and incorporation status.

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

Large table with categories: CONTRACT, REAL PROPERTY, TORTS, 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

Brief description of cause:

Defendants fraudulently advertise their decongestant products containing phenylephrine.

VII. REQUESTED IN COMPLAINT:

CHECK IF THIS IS A CLASS ACTION UNDER RULE 23, Fed. R. Civ. P. DEMAND \$ 5,000,000.00

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 09/22/2023

SIGNATURE OF ATTORNEY OF RECORD

/s/ Sarah N. Westcot