

UNITED STATES DISTRICT COURT
WESTERN DISTRICT OF WASHINGTON AT TACOMA

JEFFERSON COUNTY,

Plaintiff,

No. 3:18-cv-05661

v.

COMPLAINT

PURDUE PHAMA, L.P.; PURDUE PHARMA,
INC.; THE PURDUE FREDERICK
COMPANY, INC.; ENDO HEALTH
SOLUTIONS INC.; ENDO
PHARMACEUTICALS, INC.; JANSSEN
PHARMACEUTICALS, INC.; JOHNSON &
JOHNSON; TEVA PHARMACEUTICALS
INDUSTRIES, LTD.; TEVA
PHARMACEUTICALS USA, INC.;
CEPHALON, INC.; ALLERGAN PLC f/k/a
ACTAVIS PLC; ALLERGAN FINANCE, LLC
f/k/a ACTAVIS, INC. f/k/a WATSON
PHARMACEUTICALS, INC.; WATSON
LABORATORIES, INC.; ACTAVIS LLC;
ACTAVIS PHARMA, INC. f/k/a WATSON
PHARMA, INC; MALLINCKRODT PLC;
MALLINCKRODT, LLC; CARDINAL
HEALTH, INC.; MCKESSON
CORPORATION; AMERISOURCEBERGEN
DRUG CORPORATION; and JOHN AND
JANE DOES 1 THROUGH 100, INCLUSIVE,

JURY DEMAND

Defendants.

COMPLAINT
(3:18-cv-05661)

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TABLE OF CONTENTS

I.	INTRODUCTION	1
II.	PARTIES	6
III.	JURISDICTION AND VENUE	14
IV.	FACTUAL ALLEGATIONS	14
A.	Making an Old Drug New Again.....	14
1.	A history and background of opioids in medicine	14
2.	The Sackler family pioneered the integration of advertising and medicine.	18
3.	Purdue and the development of OxyContin.....	21
B.	The Booming Business of Addiction.....	25
1.	Other Manufacturing Defendants leapt at the opioid opportunity.	25
2.	Distributor Defendants knowingly supplied dangerous quantities of opioids while advocating for limited oversight and enforcement.	29
3.	Pill mills and overprescribing doctors also placed their financial interests ahead of their patients’ interests.	32
4.	Widespread prescription opioid use broadened the market for heroin and fentanyl.	35
C.	The Manufacturing Defendants Promoted Prescription Opioids Through Several Channels.	37
1.	The Manufacturing Defendants aggressively deployed sales representatives to push their products.	38
2.	The Manufacturing Defendants bankrolled seemingly independent “front groups” to promote opioid use and fight restrictions on opioids.	44
3.	“It was pseudoscience”: the Manufacturing Defendants paid prominent physicians to promote their products.	49

1	4.	The Manufacturing Defendants used “unbranded”	
2		advertising as a platform for their misrepresentations about	
3		opioids.....	55
4	D.	Specific Misrepresentations Made by the Manufacturing	
5		Defendants.	56
6	1.	The Manufacturing Defendants falsely claimed that the risk	
7		of opioid abuse and addiction was low.	56
8	2.	The Manufacturing Defendants falsely claimed that opioids	
9		were proven effective for chronic pain and would improve	
10		quality of life.....	69
11	3.	The Manufacturing Defendants falsely claimed doctors and	
12		patients could increase opioid usage indefinitely without	
13		added risk.	73
14	4.	The Manufacturing Defendants falsely instructed doctors	
15		and patients that more opioids were the solution when	
16		patients presented symptoms of addiction.	78
17	5.	The Manufacturing Defendants falsely claimed that risk-	
18		mitigation strategies, including tapering and abuse-	
19		deterrent technologies, made it safe to prescribe opioids for	
20		chronic use.	81
21	E.	Research by Washington State’s Department of Labor and	
22		Industries Highlights the Falseness of the Manufacturing	
23		Defendants’ Claims.....	87
24	F.	The 2016 CDC Guideline and Other Recent Studies Confirm That	
25		the Manufacturing Defendants’ Statements About the Risks and	
26		Benefits of Opioids Are Patently False.....	89
	G.	Jefferson County Has Been Directly Affected by the Opioid	
		Epidemic Caused by Defendants.	96
	H.	No Federal Agency Action, Including by the FDA, Can Provide	
		the Relief Jefferson County Seeks Here.	99
	V.	CLAIMS FOR RELIEF	100
		COUNT ONE — VIOLATIONS OF THE WASHINGTON CONSUMER	
		PROTECTION ACT, RCW 19.86, <i>ET SEQ.</i>	100
		COUNT TWO — PUBLIC NUISANCE	102

1	COUNT THREE — NEGLIGENCE.....	104
2	COUNT FOUR — GROSS NEGLIGENCE.....	105
3	COUNT FIVE — UNJUST ENRICHMENT.....	107
4	COUNT SIX — VIOLATIONS OF THE RACKETEER INFLUENCED AND	
5	CORRUPT ORGANIZATIONS ACT (“RICO”), 18 U.S.C. § 1961, <i>ET SEQ.</i>	107
6	A. Description of the Defendants’ Enterprises	108
7	B. The Enterprises Sought to Fraudulently Increase Defendants’	
8	Profits and Revenues.....	111
9	C. Predicate Acts: Mail and Wire Fraud.....	115
10	D. Jefferson County Has Been Damaged by Defendants’ RICO	
11	Violations.....	123
12	PRAYER FOR RELIEF	124
13	JURY TRIAL DEMAND	125

I. INTRODUCTION

1. The United States is experiencing the worst man-made epidemic in modern medical history—the misuse, abuse, and over-prescription of opioids.

2. Since 2000, more than 300,000 Americans have lost their lives to an opioid overdose, more than five times as many American lives as were lost in the entire Vietnam War. On any given day, 145 people will die from opioid overdoses in the United States. Drug overdoses are now the leading cause of death for Americans under age fifty.

3. The opioid crisis has become a public health emergency of unprecedented levels. Plaintiff Jefferson County, in Western Washington State and home to over 31,000 residents, has been deeply affected by the crisis. The opioid abuse prevalent throughout the County has affected Plaintiff in numerous ways, not only through the need for increased emergency medical services, but also through increased drug-related offenses affecting law enforcement, corrections, and courts, and through additional resources spent on community and social programs, including for the next generation of Jefferson County residents, who are growing up in the shadow of the opioid epidemic.

4. Jefferson County has been working to confront the epidemic caused by Defendants' reckless promotion and distribution of prescription opioids. For example, in the fall of 2016, Jefferson County, working with neighboring Kitsap and Clallam Counties as part of the Olympic Community of Health, initiated the Three-County Coordinated Opioid Response Project (3CCORP), a multi-sector collaborative effort to address the opioid epidemic in the region.

5. While the County has committed considerable resources to dealing with the crisis, fully addressing the crisis requires that those responsible for it pay for their conduct and to abate the nuisance and harms they have created in Jefferson County.

1 6. The opioid epidemic is no accident. On the contrary, it is the foreseeable
2 consequence of Defendants' reckless promotion and distribution of potent opioids for chronic
3 pain while deliberately downplaying the significant risks of addiction and overdose.

4 7. Defendant Purdue set the stage for the opioid epidemic, through the production
5 and promotion of its blockbuster drug, OxyContin. Purdue introduced a drug with a narcotic
6 payload many times higher than that of previous prescription painkillers, while executing a
7 sophisticated, multi-pronged marketing campaign to change prescribers' perception of the risk of
8 opioid addiction and to portray opioids as effective treatment for chronic pain. Purdue pushed its
9 message of opioids as a low-risk panacea on doctors and the public through every available
10 avenue, including through direct marketing, front groups, key opinion leaders, unbranded
11 advertising, and hundreds of sales representatives who visited doctors and clinics on a regular
12 basis.

13 8. As sales of OxyContin and Purdue's profits surged, Defendants Endo, Janssen,
14 Cephalon, Actavis, and Mallinckrodt—as explained in further detail below—added additional
15 prescription opioids, aggressive sales tactics, and dubious marketing claims of their own to the
16 deepening crisis. They paid hundreds of millions of dollars to market and promote the drugs,
17 notwithstanding their dangers, and pushed bought-and-paid-for “science” supporting the safety
18 and efficacy of opioids that lacked any basis in fact or reality. Obscured from the marketing was
19 the fact that prescription opioids are not much different than heroin—indeed on a molecular
20 level, they are virtually indistinguishable.

21 9. The opioid epidemic simply could not have become the crisis it is today without
22 an enormous supply of pills. Defendants McKesson, Cardinal Health, and AmerisourceBergen
23 raked in huge profits from the distribution of opioids around the United States. These companies
24 knew precisely the quantities of potent narcotics they were delivering to communities across the
25 country, including Jefferson County. Yet not only did these defendants intentionally disregard
26 their monitoring and reporting obligations under federal law, they also actively sought to evade

1 restrictions and obtain higher quotas to enable the distribution of even larger shipments of
2 opioids.

3 10. Defendants' efforts were remarkably successful: since the mid-1990s, opioids
4 have become the most prescribed class of drugs in America. Between 1991 and 2011, opioid
5 prescriptions in the U.S. tripled from 76 million to 219 million per year.¹ In 2013, health care
6 providers wrote more than 249 million prescriptions for opioid pain medication, enough for
7 every adult in the United States to have more than one bottle of pills.² In terms of annual sales,
8 the increase has been ten-fold; before the FDA approved OxyContin in 1995, annual opioid sales
9 hovered around \$1 billion. By 2015, they increased to almost \$10 billion. By 2020, revenues are
10 projected to grow to \$18 billion.³

11 11. But Defendants' profits have come at a steep price. Opioids are now the leading
12 cause of accidental death in the U.S., surpassing deaths caused by car accidents. Opioid overdose
13 deaths (which include prescription opioids as well as heroin) have risen steadily every year, from
14 approximately 8,048 in 1999, to 20,422 in 2009, to over 33,091 in 2015. In 2016, that toll
15 climbed to 42,249.⁴ As shown in the graph below, the recent surge in opioid-related deaths
16 involves prescription opioids, heroin, and other synthetic opioids. Nearly half of all opioid
17 overdose deaths involve a prescription opioid like those manufactured by Defendants,⁵ and the
18
19

20 ¹ Nora D. Volkow, MD, *America's Addiction to Opioids: Heroin and Prescription Drug Abuse*, Appearing before
21 the Senate Caucus on International Narcotics Control, NIH Nat'l Inst. on Drug Abuse (May 14, 2014),
22 [https://www.drugabuse.gov/about-nida/legislative-activities/testimony-to-congress/2016/americas-addiction-to-](https://www.drugabuse.gov/about-nida/legislative-activities/testimony-to-congress/2016/americas-addiction-to-opioids-heroin-prescription-drug-abuse)
23 [opioids-heroin-prescription-drug-abuse](https://www.drugabuse.gov/about-nida/legislative-activities/testimony-to-congress/2016/americas-addiction-to-opioids-heroin-prescription-drug-abuse).

24 ² CDC *Guideline for Prescribing Opioids for Chronic Pain*, Ctrs. for Disease Control and Prevention,
25 https://www.cdc.gov/drugoverdose/pdf/guidelines_at-a-glance-a.pdf (last visited Aug. 13, 2018).

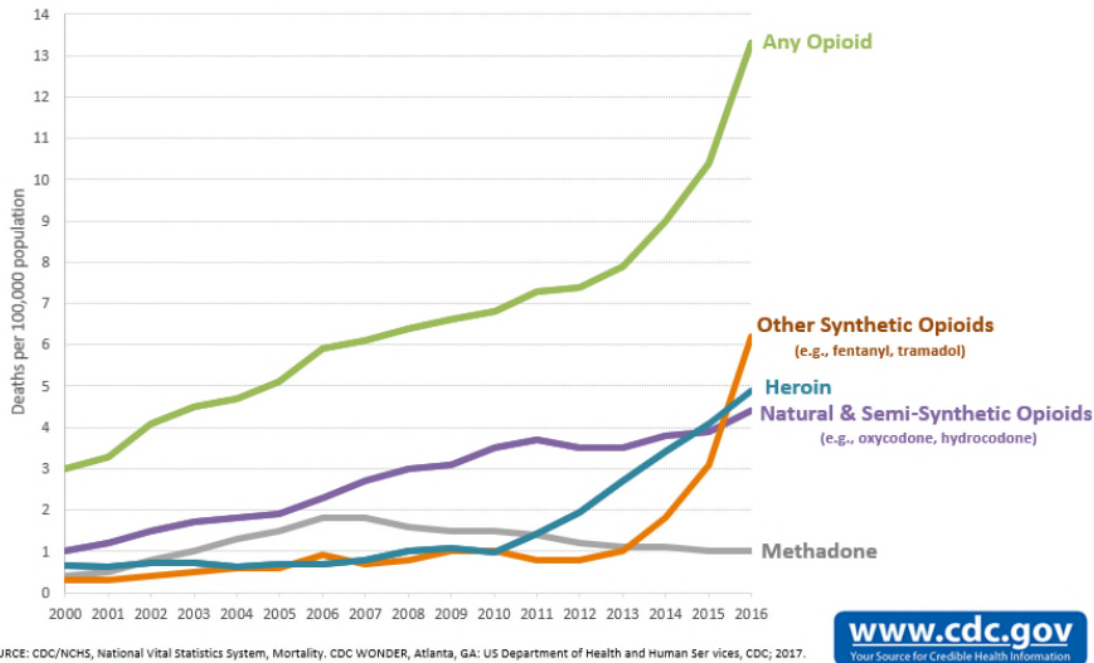
26 ³ *Report: Opioid pain sales to hit \$18.4B in the U.S. by 2020*, CenterWatch (July 17, 2017),
<https://www.centerwatch.com/news-online/2017/07/17/report-opioid-pain-sales-hit-18-4b-u-s-2020/#more-31534>.

⁴ *Overdose Death Rates*, NIH Nat'l Inst. on Drug Abuse, [https://www.drugabuse.gov/related-topics/trends-](https://www.drugabuse.gov/related-topics/trends-statistics/overdose-death-rates)
[statistics/overdose-death-rates](https://www.drugabuse.gov/related-topics/trends-statistics/overdose-death-rates) (revised Sept. 2017); *Drug Overdose Death Data*, Ctrs. for Disease Control and
Prevention, <https://www.cdc.gov/drugoverdose/data/statedeaths.html> (last updated Dec. 19, 2017).

⁵ *Understanding the Epidemic*, Ctrs. for Disease Control and Prevention,
<https://www.cdc.gov/drugoverdose/epidemic/index.html> (last updated Aug. 30, 2017).

increase in overdoses from non-prescription opioids is directly attributable to Defendants' success in expanding the market for opioids of any kind.

Overdose Deaths Involving Opioids, by Type of Opioid, United States, 2000-2016



12. To put these numbers in perspective: in 1970, when a heroin epidemic swept the U.S., there were fewer than 3,000 heroin overdose deaths. And in 1988, around the height of the crack epidemic, there were fewer than 5,000 crack overdose deaths recorded. In 2005, at its peak, methamphetamine was involved in approximately 4,500 deaths.

13. Beyond the human cost, the CDC recently estimated that the total economic burden of prescription opioid abuse costs the United States \$78.5 billion per year, which includes increased costs for health care and addiction treatment, increased strains on human services and criminal justice systems, and substantial losses in workforce productivity.⁶

14. But even these estimates are conservative. The Council of Economic Advisers—the primary advisor to the Executive Office of the President—recently issued a report estimating

⁶ CDC Foundation's *New Business Pulse Focuses on Opioid Overdose Epidemic*, Ctrs. for Disease Control and Prevention (Mar. 15, 2017), <https://www.cdc.gov/media/releases/2017/a0315-business-pulse-opioids.html>.

1 that “in 2015, the economic cost of the opioid crisis was \$504.0 billion, or 2.8% of GDP that
2 year. This is over six times larger than the most recently estimated economic cost of the
3 epidemic.”⁷ Whatever the final tally, there is no doubt that this crisis has had a profound
4 economic impact.

5 15. Defendants orchestrated this crisis. Despite knowing about the true hazards of
6 their products, Defendants misleadingly advertised their opioids as safe and effective for treating
7 chronic pain and pushed hundreds of millions of pills into the marketplace for consumption.
8 Through their sophisticated and well-orchestrated campaign, Defendants touted the purported
9 benefits of opioids to treat pain and downplayed the risks of addiction. Moreover, even as the
10 deadly toll of prescription opioid use became apparent to Defendants in years following
11 OxyContin’s launch, Defendants persisted in aggressively selling and distributing prescription
12 opioids, while evading their monitoring and reporting obligations, so that massive quantities of
13 addictive opioids continued to pour into Jefferson County and other communities around the
14 United States.

15 16. Defendants consistently, deliberately, and recklessly made and continue to make
16 false and misleading statements regarding, among other things, the low risk of addiction to
17 opioids, opioids’ efficacy for chronic pain and ability to improve patients’ quality of life with
18 long-term use, the lack of risk associated with higher dosages of opioids, the need to prescribe
19 more opioids to treat withdrawal symptoms, and that risk-mitigation strategies and abuse-
20 deterrent technologies allow doctors to safely prescribe opioids.

21 17. Because of Defendants’ misconduct, Jefferson County is experiencing a severe
22 public health crisis and has suffered significant economic damages, including but not limited to
23 increased costs related to public health, opioid-related crimes and emergencies, criminal justice,
24

25 ⁷ *The Underestimated Cost of the Opioid Crisis*, The Council of Econ. Advisers (Nov. 2017),
26 <https://static.politico.com/1d/33/4822776641cfbac67f9bc7dbd9c8/the-underestimated-cost-of-the-opioid-crisis-embargoed.pdf>.

1 and public safety. Jefferson County has incurred substantial costs in responding to the crisis and
2 will continue to do so in the future.

3 18. Accordingly, Jefferson County brings this action to hold Defendants liable for
4 their misrepresentations regarding the benefits and risks of opioids, as well as for their failure to
5 monitor, detect, investigate, and report suspicious orders of prescription opioids. This conduct (i)
6 violates the Washington Consumer Protection Act, RCW 19.86 *et seq.*, (ii) constitutes a public
7 nuisance under Washington law, (iii) constitutes negligence and gross negligence under
8 Washington law, (iv) has unjustly enriched Defendants, and (v) violates the Racketeer Influenced
9 and Corrupt Organizations Act (“RICO”), 18 U.S.C. §1961, *et seq.*

10 II. PARTIES

11 Jefferson County

12 19. Plaintiff Jefferson County (“Plaintiff” or “Jefferson County” or “County”) is a
13 Washington County organized and existing under the laws of the State of Washington, RCW
14 36.01 *et seq.*

15 Purdue

16 20. Defendant Purdue Pharma, L.P. is a limited partnership organized under the laws
17 of Delaware. Defendant Purdue Pharma, Inc. is a New York corporation with its principal place
18 of business in Stamford, Connecticut. Defendant The Purdue Frederick Company is a Delaware
19 corporation with its principal place of business in Stamford, Connecticut. Collectively, these
20 entities are referred to as “Purdue.”

21 21. Each Purdue entity acted in concert with one another and acted as agents and/or
22 principals of one another in connection with the conduct described herein.

23 22. Purdue manufactures, promotes, sells, markets, and distributes opioids such as
24 OxyContin, MS Contin, Dilaudid/Dilaudid HP, Butrans, Hysingla ER, and Targiniq ER.

23. Purdue generates substantial sales revenue from its opioids. For example, OxyContin is Purdue's best-selling opioid, and since 2009, Purdue has generated between \$2 and \$3 billion annually in sales of OxyContin alone.

Endo

24. Defendant Endo Pharmaceuticals, Inc. is a wholly owned subsidiary of Defendant Endo Health Solutions Inc. Both are Delaware corporations with their principal place of business in Malvern, Pennsylvania. Collectively, these entities are referred to as "Endo."

25. Each Endo entity acted in concert with one another and acted as agents and/or principals of one another in connection with the conduct described herein.

26. Endo manufactures, promotes, sells, markets, and distributes opioids such as Percocet, Opana, and Opana ER.

27. Endo generates substantial sales from its opioids. For example, opioids accounted for more than \$400 million of Endo's overall revenues of \$3 billion in 2012, and Opana ER generated more than \$1 billion in revenue for Endo in 2010 and 2013.

Janssen and Johnson & Johnson

28. Defendant Janssen Pharmaceuticals, Inc. is a Pennsylvania corporation with its principal place of business in Titusville, New Jersey, and is a wholly owned subsidiary of Defendant Johnson & Johnson, a New Jersey corporation with its principal place of business in New Brunswick, New Jersey. Collectively, these entities are referred to as "Janssen."

29. Both entities above acted in concert with one another and acted as agents and/or principals of one another in connection with the conduct described herein.

30. Johnson & Johnson is the only company that owns more than 10% of Janssen Pharmaceuticals, Inc., and corresponds with the FDA regarding the drugs manufactured by Janssen Pharmaceuticals, Inc. Johnson & Johnson also paid prescribers to speak about opioids manufactured by Janssen Pharmaceuticals, Inc. In short, Johnson & Johnson controls the sale and development of the drugs manufactured by Janssen Pharmaceuticals, Inc.

31. Janssen manufactures, promotes, sells, markets, and distributes opioids such as Duragesic, Nucynta, and Nucynta ER. Janssen stopped manufacturing Nucynta and Nucynta ER in 2015.

32. Janssen generates substantial sales revenue from its opioids. For example, Duragesic accounted for more than \$1 billion in sales in 2009, and Nucynta and Nucynta ER accounted for \$172 million in sales in 2014.

Cephalon and Teva

33. Defendant Cephalon, Inc. (“Cephalon”) is a Delaware corporation with its principal place of business in Frazer, Pennsylvania. Defendant Teva Pharmaceutical Industries, Ltd. (“Teva Ltd.”) is an Israeli corporation with its principal place of business in Petah Tikva, Israel. In 2011, Teva Ltd. acquired Cephalon. Defendant Teva Pharmaceuticals USA, Inc. (“Teva USA”) is a Delaware corporation and a wholly owned subsidiary of Teva Ltd. in Pennsylvania. Teva USA acquired Cephalon in October 2011.

34. Cephalon manufactures, promotes, sells, and distributes opioids, including Actiq and Fentora, in the United States.

35. Teva Ltd., Teva USA, and Cephalon work together closely to market and sell Cephalon products in the United States. Teva Ltd. conducts all sales and marketing activities for Cephalon in the United States through Teva USA and has done so since its October 2011 acquisition of Cephalon. Teva Ltd. and Teva USA hold out Actiq and Fentora as Teva products to the public. Teva USA sells all former Cephalon-branded products through its “specialty medicines” division. The FDA-approved prescribing information and medication guide, which are distributed with Cephalon opioids, disclose that the guide was submitted by Teva USA, and directs physicians to contact Teva USA to report adverse events.

36. All of Cephalon’s promotional websites, including those for Actiq and Fentora, display Teva Ltd.’s logo.⁸ Teva Ltd.’s financial reports list Cephalon’s and Teva USA’s sales as

⁸ Actiq, <http://www.actiq.com/> (last visited Aug. 13, 2018).

its own, and its year-end report for 2012—the year following the Cephalon acquisition in October 2011—attributed a 22% increase in its specialty medicine sales to “the inclusion of a full year of Cephalon’s specialty sales,” including sales of Fentora.⁹ Through interrelated operations like these, Teva Ltd. operates in the United States through its subsidiaries Cephalon and Teva USA. The United States is the largest of Teva Ltd.’s global markets, representing 53% of its global revenue in 2015, and, were it not for the existence of Teva USA and Cephalon, Teva Ltd. would conduct those companies’ business in the United States itself.

37. Upon information and belief, Teva Ltd. directs the business practices of Cephalon and Teva USA, and their profits inure to the benefit of Teva Ltd. as controlling shareholder. Collectively, these entities are referred to as “Cephalon.”

Allergan, Actavis, and Watson

38. Defendant Allergan plc is a public limited company incorporated in Ireland with its principal place of business in Dublin, Ireland. Actavis plc acquired Allergan, Inc. in March 2015, and the combined company changed its name to Allergan plc in June 2015. Actavis plc (formerly known as Actavis Limited) was incorporated in Ireland in May 2013 for the merger between Actavis, Inc. and Warner Chilcott plc.

39. Defendant Watson Pharmaceuticals, Inc. acquired Actavis Group in October 2012 and changed its name to Actavis, Inc. as of January 2013.

40. Defendant Allergan Finance, LLC (formerly known as Actavis, Inc.) is based in Parsippany, New Jersey. It operates as a subsidiary of Allergan plc.

41. Defendant Watson Laboratories, Inc. is a Nevada corporation with its principal place of business in Corona, California, and is a wholly owned subsidiary of Allergan plc (f/k/a Actavis, Inc., f/k/a Watson Pharmaceuticals, Inc.).

⁹ *Teva Pharm. Indus. Ltd. Form 20-F*, U.S. Sec. and Exchange Commission (Feb. 12, 2013), http://annualreports.com/HostedData/AnnualReportArchive/t/NASDAQ_TEVA_2012.pdf.

42. Defendant Actavis Pharma, Inc. is registered to do business with the Washington Secretary of State as a Delaware corporation with its principal place of business in New Jersey and was formerly known as Watson Pharma, Inc.

43. Defendant Actavis LLC is a Delaware limited liability company with its principal place of business in Parsippany, New Jersey.

44. Each of these defendants and entities is owned by Defendant Allergan plc, which uses them to market and sell its drugs in the United States. Upon information and belief, Defendant Allergan plc exercises control over these marketing and sales efforts and profits from the sale of Allergan/Actavis/Watson products ultimately inure to its benefit. Collectively, these defendants and entities are referred to as "Actavis."

45. Actavis manufactures, promotes, sells, and distributes opioids, including the branded drugs Kadian and Norco and generic versions of Kadian, Duragesic, and Opana in the United States. Actavis acquired the rights to Kadian from King Pharmaceuticals, Inc. on December 30, 2008, and began marketing Kadian in 2009.

Mallinckrodt

46. Defendant Mallinckrodt plc is an Irish public limited company headquartered in Staines-upon-Thames, United Kingdom, with its U.S. headquarters in St. Louis, Missouri. Mallinckrodt plc was incorporated in January 2013 for the purpose of holding the pharmaceuticals business of Covidien plc, which was fully transferred to Mallinckrodt in June of that year. Mallinckrodt began as a U.S.-based company, with the founding of Mallinckrodt & Co. in 1867; Tyco International Ltd. acquired the company in 2000. In 2008, Tyco Healthcare Group separated from Tyco International and renamed itself Covidien.

47. Defendant Mallinckrodt, LLC is a limited liability company organized and existing under the laws of the State of Delaware and licensed to do business in Washington. Mallinckrodt, LLC is a wholly owned subsidiary of Mallinckrodt plc. Mallinckrodt plc and Mallinckrodt, LLC are referred to as "Mallinckrodt."

1 48. Mallinckrodt manufactures, markets, and sells drugs in the United States. As of
2 2012, it was the largest U.S. supplier of opioid pain medications. In particular, it is one of the
3 largest manufacturers of oxycodone in the U.S.

4 49. Mallinckrodt currently manufactures and markets two branded opioids: Exalgo,
5 which is extended-release hydromorphone, sold in 8, 12, 16, and 32 mg dosage strengths, and
6 Roxicodone, which is oxycodone, sold in 15 and 30 mg dosage strengths. In addition,
7 Mallinckrodt previously developed, promoted, and sold the following branded opioid products:
8 Magnacet, TussiCaps, and Xartemis XR.

9 50. While it has sought to develop its branded opioid products, Mallinckrodt has long
10 been a leading manufacturer of generic opioids. Mallinckrodt estimated that in 2015 it received
11 approximately 25% of the U.S. Drug Enforcement Administration's ("DEA") entire annual quota
12 for controlled substances that it manufactures. Mallinckrodt also estimated, based on IMS Health
13 data for the same period, that its generics claimed an approximately 23% market share of DEA
14 Schedules II and III opioid and oral solid dose medications.

15 51. Mallinckrodt operates a vertically integrated business in the United States: (1)
16 importing raw opioid materials, (2) manufacturing generic opioid products, primarily at its
17 facility in Hobart, New York, and (3) marketing and selling its products to drug distributors,
18 specialty pharmaceutical distributors, retail pharmacy chains, pharmaceutical benefit managers
19 that have mail-order pharmacies, and hospital buying groups.

20 52. In 2017, Mallinckrodt agreed to settle for \$35 million the Department of Justice's
21 allegations regarding excessive sales of oxycodone in Florida. The Department of Justice alleged
22 that even though Mallinckrodt knew that its oxycodone was being diverted to illicit use, it
23 nonetheless continued to incentivize and supply these suspicious sales, and it failed to notify the
24 DEA of the suspicious orders in violation of its obligations as a registrant under the Controlled
25 Substances Act, 21 U.S.C. § 801 *et seq.* ("CSA").
26

53. Defendants Purdue, Endo, Janssen, Cephalon, Actavis, and Mallinckrodt are collectively referred to as the “Manufacturing Defendants.”

AmerisourceBergen

54. Defendant AmerisourceBergen Drug Corporation (“AmerisourceBergen”) is a Delaware corporation with its principal place of business located in Chesterbrook, Pennsylvania.

55. According to its 2016 Annual Report, AmerisourceBergen is “one of the largest global pharmaceutical sourcing and distribution services companies” with “over \$145 billion in annual revenue.”

56. AmerisourceBergen is licensed as a “wholesale distributor” to sell prescription and non-prescription drugs in Washington State, including opioids. It operates a warehouse in Kent, Washington.

Cardinal Health

57. Defendant Cardinal Health, Inc. (“Cardinal Health”) is an Ohio Corporation with its principal place of business in Dublin, Ohio.

58. According to its 2017 Annual Report, Cardinal Health is “a global, integrated healthcare services and products company serving hospitals, healthcare systems, pharmacies, ambulatory surgery centers, clinical laboratories and physician offices worldwide . . . deliver[ing] medical products and pharmaceuticals.” In 2017 alone, Cardinal Health generated revenues of nearly \$130 billion.

59. Cardinal Health is licensed as a “wholesale distributor” to sell prescription and non-prescription drugs in Washington State, including opioids. It operates a warehouse in Fife, Washington.

McKesson

60. Defendant McKesson Corporation (“McKesson”) is a Delaware Corporation with its principal place of business in San Francisco, California.

61. McKesson is the largest pharmaceutical distributor in North America, delivering nearly one-third of all pharmaceuticals used in this region.

62. According to its 2017 Annual Report, McKesson “partner[s] with pharmaceutical manufacturers, providers, pharmacies, governments and other organizations in healthcare to help provide the right medicines, medical products and healthcare services to the right patients at the right time, safely and cost-effectively.” Additionally, McKesson’s pharmaceutical distribution business operates and serves thousands of customer locations through a network of twenty-seven distribution centers, as well as a primary redistribution center, two strategic redistribution centers and two repackaging facilities, serving all fifty states and Puerto Rico.

63. For the fiscal year ending March 31, 2017, McKesson generated revenues of \$198.5 billion.

64. McKesson is licensed as a “wholesale distributor” to sell prescription and non-prescription drugs in Washington State, including opioids. It operates warehouses in Everett and Auburn, Washington.

65. Collectively, McKesson, AmerisourceBergen, and Cardinal Health (together “Distributor Defendants”) account for approximately 85% of all drug shipments in the United States.

John and Jane Does 1-100, inclusive

66. In addition to the Defendants identified herein, the true names, roles, and/or capacities in the wrongdoing alleged herein of Defendants named John and Jane Does 1 through 100, inclusive, are currently unknown to Plaintiff, and thus, are named as Defendants under fictitious names as permitted by the rules of this Court. Plaintiff will amend this complaint and identify their true identities and their involvement in the wrongdoing at issue, as well as the specific causes of action asserted against them when they become known.

III. JURISDICTION AND VENUE

67. This Court has jurisdiction over this action pursuant to 28 U.S.C. § 1332. The Court also has federal question subject matter jurisdiction arising out of Plaintiff's RICO claims pursuant to 28 U.S.C. § 1331 and 18 U.S.C. § 1961, *et seq.*

68. Venue in this Court is proper under 28 U.S.C. § 1391(b).

IV. FACTUAL ALLEGATIONS

A. Making an Old Drug New Again

1. A history and background of opioids in medicine

69. The term "opioid" refers to a class of drugs that bind with opioid receptors in the brain and includes natural, synthetic, and semi-synthetic opioids.¹⁰ Generally used to treat pain, opioids produce multiple effects on the human body, the most significant of which are analgesia, euphoria, and respiratory depression. In addition, opioids cause sedation and constipation.

70. Most of these effects are medically useful in certain situations, but respiratory depression is the primary limiting factor for the use of opioids. While the body develops tolerance to the analgesic and euphoric effects of opioids relatively quickly, this is not true with respect to respiratory depression. At high doses, opioids can and often do arrest respiration altogether. This is why the risk of opioid overdose is so high, and why many of those who overdose simply go to sleep and never wake up.

71. Natural opioids are derived from the opium poppy and have been used since antiquity, going as far back as 3400 B.C. The opium poppy contains various opium alkaloids, three of which are used commercially today: morphine, codeine, and thebaine.

72. A 16th-century European alchemist, Paracelsus, is generally credited with developing a tincture of opium and alcohol called laudanum, but it was a British physician a

¹⁰ At one time, the term "opiate" was used for natural opioids, while "opioid" referred to synthetic substances manufactured to mimic opiates. Now, however, most medical professionals use "opioid" to refer broadly to natural, semi-synthetic, and synthetic opioids. A fourth class of opioids, endogenous opioids (e.g., endorphins), is produced naturally by the human body.

1 century later who popularized the use of laudanum in Western medicine. “Sydenham’s
2 laudanum” was a simpler tincture than Paracelsus’s and was widely adopted as a treatment not
3 only for pain, but for coughs, dysentery, and numerous other ailments. Laudanum contains
4 almost all of the opioid alkaloids and is still available by prescription today.

5 73. Chemists first isolated the morphine and codeine alkaloids in the early 1800s, and
6 the pharmaceutical company Merck began large-scale production and commercial marketing of
7 morphine in 1827. During the American Civil War, field medics commonly used morphine,
8 laudanum, and opium pills to treat the wounded, and many veterans were left with morphine
9 addictions. It was upper and middle class white women, however, who comprised the majority of
10 opioid addicts in the late 19th-century United States, using opioid preparations widely available
11 in pain elixirs, cough suppressants, and patent medicines. By 1900, an estimated 300,000 people
12 were addicted to opioids in the United States,¹¹ and many doctors prescribed opioids solely to
13 prevent their patients from suffering withdrawal symptoms.

14 74. Trying to develop a drug that could deliver opioids’ potent pain relief without
15 their addictive properties, chemists continued to isolate and refine opioid alkaloids. Heroin, first
16 synthesized from morphine in 1874, was marketed commercially by the Bayer Pharmaceutical
17 Company beginning in 1898 as a safe alternative to morphine. Heroin’s market position as a safe
18 alternative was short-lived, however; Bayer stopped mass-producing heroin in 1913 because of
19 its dangers. German chemists then looked to the alkaloid thebaine, synthesizing oxymorphone
20 and oxycodone from thebaine in 1914 and 1916, respectively, with the hope that the different
21 alkaloid source might provide the benefits of morphine and heroin without the drawbacks.

22 75. But each opioid was just as addictive as the one before it, and eventually the issue
23 of opioid addiction could not be ignored. The nation’s first Opium Commissioner, Hamilton
24 Wright, remarked in 1911, “The habit has this nation in its grip to an astonishing extent. Our

25 ¹¹ Nick Miroff, *From Teddy Roosevelt to Trump: How drug companies triggered an opioid crisis a century ago*,
26 Washington Post (Oct. 17, 2017), https://www.washingtonpost.com/news/retropolis/wp/2017/09/29/the-greatest-drug-fiends-in-the-world-an-american-opioid-crisis-in-1908/?utm_term=.7832633fd7ca.

1 prisons and our hospitals are full of victims of it, it has robbed ten thousand businessmen of
2 moral sense and made them beasts who prey upon their fellows . . . it has become one of the
3 most fertile causes of unhappiness and sin in the United States.”¹²

4 76. Concerns over opioid addiction led to national legislation and international
5 agreements regulating narcotics: the International Opium Convention, signed at the Hague in
6 1912, and, in the U.S., the Harrison Narcotics Tax Act of 1914. Opioids were no longer marketed
7 as cure-alls and instead were relegated to the treatment of acute pain.

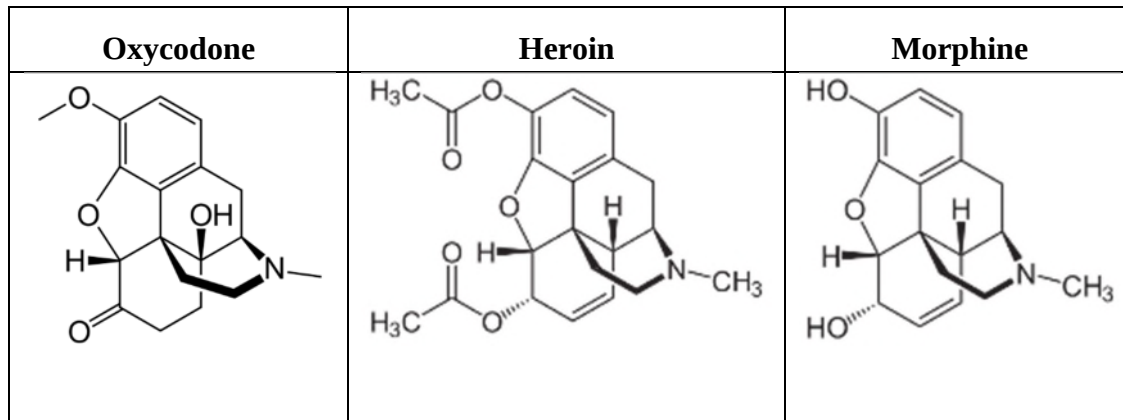
8 77. Throughout the twentieth century, pharmaceutical companies continued to
9 develop prescription opioids, but these opioids were generally produced in combination with
10 other drugs, with relatively low opioid content. For example, Percodan, produced by Defendant
11 Endo since 1950, is oxycodone and aspirin, and contains just under 5 mg of oxycodone.
12 Percocet, manufactured by Endo since 1971, is the combination of oxycodone and
13 acetaminophen, with dosage strengths delivering between 2.5 mg and 10 mg of oxycodone.
14 Vicodin, a combination of hydrocodone and acetaminophen, was introduced in the U.S. in 1978
15 and is sold in strengths of 5 mg, 7.5 mg, and 10 mg of hydrocodone. Defendant Janssen also
16 manufactured a drug with 5 mg of oxycodone and 500 mg of acetaminophen, called Tylox, from
17 1984 to 2012.

18 78. In contrast, OxyContin, the product with the dubious honor of the starring role in
19 the opioid epidemic, is pure oxycodone. Purdue initially made it available in the following
20 dosage strengths: 10 mg, 15 mg, 20 mg, 30 mg, 40 mg, 60 mg, 80 mg, and 160 mg. In other
21 words, the weakest OxyContin delivers as much narcotic as the strongest Percocet, and some
22 OxyContin tablets delivered sixteen times as much as that.

23 79. Prescription opioids are essentially pharmaceutical heroin; they are synthesized
24 from the same plant, have similar molecular structures, and bind to the same receptors in the
25 human brain. It is no wonder then that there is a straight line between prescription opioid abuse

26 ¹² *Id.*

and heroin addiction. Indeed, studies show that over 80% of new heroin addicts between 2008 and 2010 started with prescription opioids.¹³



80. Medical professionals describe the strength of various opioids in terms of “morphine milligram equivalents” (“MME”). According to the CDC, dosages at or above 50 MME/day double the risk of overdose compared to 20 MME/day, and one study found that patients who died of opioid overdose were prescribed an average of 98 MME/day.

81. Different opioids provide varying levels of MMEs. For example, just 33 mg of oxycodone provides 50 MME. Thus, at OxyContin’s twice-daily dosing, the 50 MME/day threshold is reached by a prescription of 15 mg twice daily. One 160 mg tablet of OxyContin, which Purdue took off the market in 2001, delivered 240 MME.¹⁴

82. As journalist Barry Meier wrote in his 2003 book *Pain Killer: A “Wonder” Drug’s Trail of Addiction and Death*, “In terms of narcotic firepower, OxyContin was a nuclear weapon.”¹⁵

¹³ Jones CM, *Heroin use and heroin use risk behaviors among nonmedical users of prescription opioid pain relievers - United States, 2002-2004 and 2008-2010*, 132(1-2) Drug Alcohol Depend. 95-100 (Sept. 1, 2013), <https://www.ncbi.nlm.nih.gov/pubmed/23410617>.

¹⁴ The wide variation in the MME strength of prescription opioids renders misleading any effort to capture “market share” by the number of pills or prescriptions attributed to Purdue or other manufacturers. Purdue, in particular, focuses its business on branded, highly potent pills, causing it to be responsible for a significant percent of the total amount of MME in circulation even though it currently claims to have a small percent of the market share in terms of pills or prescriptions.

¹⁵ Barry Meier, *Pain Killer: A “Wonder” Drug’s Trail of Addiction and Death* (Rodale 2003).

83. Fentanyl, an even more potent and more recent arrival in the opioid tale, is a synthetic opioid that is 100 times stronger than morphine and 50 times stronger than heroin. First developed in 1959 by Dr. Paul Janssen under a patent held by Janssen Pharmaceutica, fentanyl is increasingly prevalent in the market for opioids created by Defendants' promotion, with particularly lethal consequences. In many instances, illicit fentanyl is manufactured to look like oxycodone tablets, in the light blue color and with the "M" stamp of Defendant Mallinckrodt's 30mg oxycodone pills. These lookalike pills have been found around the country, including in Washington State.¹⁶

2. The Sackler family pioneered the integration of advertising and medicine.

84. Given the history of opioid use in the U.S. and the medical profession's resulting wariness, the commercial success of Defendants' prescription opioids would not have been possible without a fundamental shift in prescribers' perception of the risks and benefits of long-term opioid use.

85. As it turned out, Purdue was uniquely positioned to execute just such a maneuver, thanks to the legacy of a man named Arthur Sackler. The Sackler family is the sole owner of Purdue and one of the wealthiest families in America, surpassing the wealth of storied families like the Rockefellers, the Mellons, and the Busches.¹⁷ Because of Purdue and, in particular, OxyContin, the Sacklers' net worth was \$13 billion as of 2016. Today, all nine members of the Purdue board are family members, and all of the company's profits go to Sackler family trusts and entities.¹⁸ Yet the Sacklers have avoided publicly associating themselves with Purdue, letting others serve as the spokespeople for the company.

¹⁶ See e.g., Sharon Bogan, *Illicit fentanyl found locally in fake opioid pills*, Public Health Insider (Oct. 2, 2017), <https://publichealthinsider.com/2017/10/02/illicit-fentanyl-found-locally-in-fake-opioid-pills/>; *Mislabeled painkillers "a fatal overdose waiting to happen,"* CBS News (Feb. 29, 2016, 10:46am), <https://www.cbsnews.com/news/mislabeled-painkillers-a-fatal-overdose-waiting-to-happen/>.

¹⁷ Alex Morrell, *The OxyContin Clan: The \$14 Billion Newcomer to Forbes 2015 List of Richest U.S. Families*, Forbes (July 1, 2015, 10:17am), <https://www.forbes.com/sites/alexmorrell/2015/07/01/the-oxycontin-clan-the-14-billion-newcomer-to-forbes-2015-list-of-richest-u-s-families/#382ab3275e02>.

¹⁸ David Armstrong, *The man at the center of the secret OxyContin files*, Stat News (May 12, 2016), <https://www.statnews.com/2016/05/12/man-center-secret-oxycontin-files/>.

86. The Sackler brothers—Arthur, Mortimer, and Raymond—purchased a small patent-medicine company called The Purdue Frederick Company in 1952. While all three brothers were accomplished psychiatrists, it was Arthur, the oldest, who directed the Sackler story, treating his brothers more as his protégés than colleagues, putting them both through medical school and essentially dictating their paths. It was Arthur who created the Sackler family’s wealth, and it was Arthur who created the pharmaceutical advertising industry as we know it—laying the groundwork for the OxyContin promotion that would make the Sacklers billionaires.

87. Arthur Sackler was both a psychiatrist and a marketing executive, and, by many accounts, a brilliant and driven man. He pursued two careers simultaneously, as a psychiatrist at Creedmoor State Hospital in New York and the president of an advertising agency called William Douglas McAdams. Arthur pioneered both print advertising in medical journals and promotion through physician “education” in the form of seminars and continuing medical education courses. He understood intuitively the persuasive power of recommendations from fellow physicians, and did not hesitate to manipulate information when necessary. For example, one promotional brochure produced by his firm for Pfizer showed business cards of physicians from various cities as if they were testimonials for the drug, but when a journalist tried to contact these doctors, he discovered that they did not exist.¹⁹

88. It was Arthur who, in the 1960s, made Valium into the first \$100-million drug, so popular it became known as “Mother’s Little Helper.” His expertise as a psychiatrist was key to his success; as his biography in the Medical Advertising Hall of Fame notes, it “enabled him to position different indications for Roche’s Librium and Valium—to distinguish for the physician the complexities of anxiety and psychic tension.”²⁰ When Arthur’s client, Roche, developed Valium, it already had a similar drug, Librium, another benzodiazepine, on the market for

¹⁹ Meier, *supra* note 15, at 204.

²⁰ MAHF Inductees, Arthur M. Sackler, Med. Advert. Hall of Fame, <https://www.mahf.com/mahf-inductees/> (last visited Aug. 13, 2018).

1 treatment of anxiety. So Arthur invented a condition he called “psychic tension”—essentially
 2 stress—and pitched Valium as the solution.²¹ The campaign, for which Arthur was compensated
 3 based on volume of pills sold,²² was a remarkable success.

4 89. Arthur’s entrepreneurial drive led him to create not only the advertising for his
 5 clients but also the vehicle to bring their advertisements to doctors—a biweekly newspaper
 6 called the *Medical Tribune*, which he distributed for free to doctors nationwide. Arthur also
 7 conceived a company now called IMS Health Holdings Inc., which monitors prescribing
 8 practices of every doctor in the U.S. and sells this valuable data to pharmaceutical companies
 9 like Defendants, who utilize it to tailor their sales pitches to individual physicians.

10 90. Even as he expanded his business dealings, Arthur was adept at hiding his
 11 involvement in them. When, during a 1962 Senate hearing about deceptive pharmaceutical
 12 advertising, he was asked about a public relations company called Medical and Science
 13 Communications Associates, which distributed marketing from drug companies disguised as
 14 news articles, Arthur was able to truthfully testify that he never was an officer for nor had any
 15 stock in that company. But the company’s sole shareholder was his then-wife. Around the same
 16 time, Arthur also successfully evaded an investigative journalist’s attempt to link the Sacklers to
 17 a company called MD Publications, which had funneled payments from drug companies to an
 18 FDA official named Henry Welch, who was forced to resign when the scandal broke.²³ Arthur
 19 had set up such an opaque and layered business structure that his connection to MD Publications
 20 was only revealed decades later when his heirs were fighting over his estate.

21 91. Arthur Sackler did not hesitate to manipulate information to his advantage. His
 22 legacy is a corporate culture that prioritizes profits over people. In fact, in 2007, federal
 23 prosecutors conducting a criminal investigation of Purdue’s fraudulent advertising of OxyContin

24
 25 ²¹ Meier, *supra* note 15, at 202; *One Family Reaped Billions From Opioids*, WBUR On Point (Oct. 23, 2017),
<http://www.wbur.org/onpoint/2017/10/23/one-family-reaped-billions-from-opioids>.

26 ²² WBUR On Point interview, *supra* note 21.

²³ Meier, *supra* note 15, at 210-14.

found a “corporate culture that allowed this product to be misbranded with the intent to defraud and mislead.”²⁴ Court documents from the prosecution state that “certain Purdue supervisors and employees, with the intent to defraud or mislead, marketed and promoted OxyContin as less addictive, less subject to abuse and diversion, and less likely to cause tolerance and withdrawal than other pain medications”²⁵ Half a century after Arthur Sackler wedded advertising and medicine, Purdue employees were following his playbook, putting product sales over patient safety.

3. Purdue and the development of OxyContin

92. After the Sackler brothers acquired The Purdue Frederick Company in 1952, Purdue sold products ranging from earwax remover to antiseptic, and it became a profitable business. As an advertising executive, Arthur Sackler was not involved, on paper at least, in running Purdue because that would have been a conflict of interest. Raymond Sackler became Purdue’s head executive while Mortimer Sackler ran Purdue’s UK affiliate.

93. In the 1980s, Purdue, through its UK affiliate, acquired a Scottish drug producer that had developed a sustained-release technology suitable for morphine. Purdue marketed this extended-release morphine as MS Contin. It quickly became Purdue’s best seller. As the patent expiration for MS Contin loomed, Purdue searched for a drug to replace it. Around that time, Raymond Sackler’s oldest son, Richard Sackler, who was also a trained physician, became more involved in the management of the company. Richard Sackler had grand ambitions for the company; according to a long-time Purdue sales representative, “Richard really wanted Purdue to be big—I mean *really* big.”²⁶ Richard Sackler believed Purdue should develop another use for its “Contin” timed-release system.

²⁴ Naomi Spencer, *OxyContin manufacturer reaches \$600 million plea deal over false marketing practices*, World Socialist Web Site (May 19, 2007), <http://www.wsws.org/en/articles/2007/05/oxy-m19.html>.

²⁵ Agreed Statement of Facts, *United States v. Purdue Frederick Co.*, No. 1:07-cr-00029 (W.D. Va. May 10, 2007).

²⁶ Christopher Glazek, *The Secretive Family Making Billions from the Opioid Crisis*, Esquire (Oct. 16, 2017), <http://www.esquire.com/news-politics/a12775932/sackler-family-oxycontin/>.

94. In 1990, Purdue's VP of clinical research, Robert Kaiko, sent a memo to Richard Sackler and other executives recommending that the company work on a pill containing oxycodone. At the time, oxycodone was perceived as less potent than morphine, largely because it was most commonly prescribed as Percocet, the relatively weak oxycodone-acetaminophen combination pill. MS Contin was not only approaching patent expiration but had always been limited by the stigma associated with morphine. Oxycodone did not have that problem, and what's more, it was sometimes mistakenly called "oxycodine," which also contributed to the perception of relatively lower potency, because codeine is weaker than morphine. Purdue acknowledged using this to its advantage when it eventually pled guilty to criminal charges of "misbranding" in 2007, admitting that it was "well aware of the incorrect view held by many physicians that oxycodone was weaker than morphine" and "did not want to do anything 'to make physicians think that oxycodone was stronger or equal to morphine' or to 'take any steps . . . that would affect the unique position that OxyContin'" held among physicians.²⁷

95. For Purdue and OxyContin to be "*really big*," Purdue needed to both distance its new product from the traditional view of narcotic addiction risk, and broaden the drug's uses beyond cancer pain and hospice care. A marketing memo sent to Purdue's top sales executives in March 1995 recommended that if Purdue could show that the risk of abuse was lower with OxyContin than with traditional immediate-release narcotics, sales would increase.²⁸ As discussed below, Purdue did not find or generate any such evidence, but this did not stop Purdue from making that claim regardless.

96. Despite the fact that there has been little or no change in the amount of pain reported in the U.S. over the last twenty years, Purdue recognized an enormous untapped market for its new drug. As Dr. David Haddox, a Senior Medical Director at Purdue, declared on the Early Show, a CBS morning talk program, "There are 50 million patients in this country who

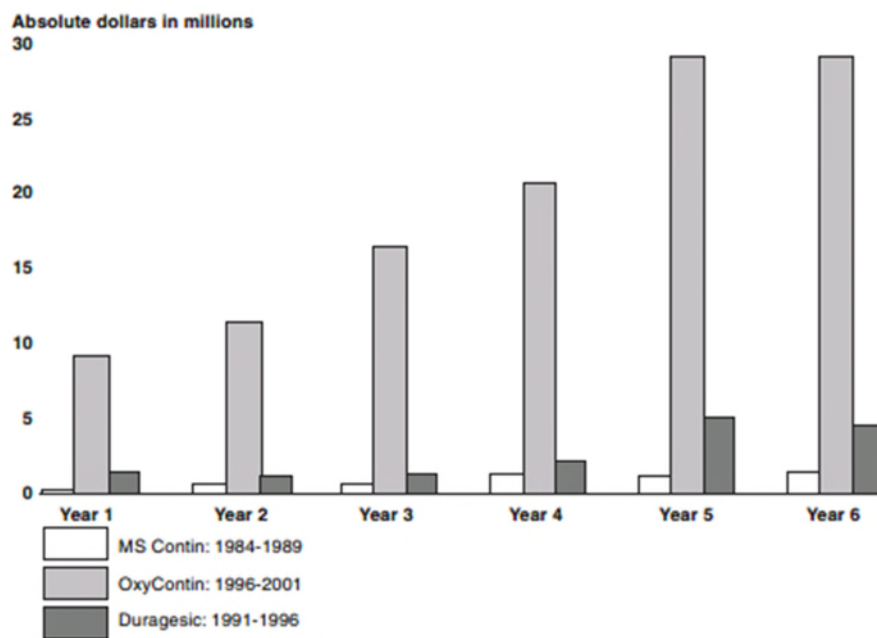
²⁷ *United States v. Purdue Frederick Co.*, *supra* note 25.

²⁸ Meier, *supra* note 15, at 269.

1 have chronic pain that's not being managed appropriately every single day. OxyContin is one of
 2 the choices that doctors have available to them to treat that."²⁹

3 97. In pursuit of these 50 million potential customers, Purdue poured resources into
 4 OxyContin's sales force and advertising. The graph below shows how promotional spending in
 5 the first six years following OxyContin's launch dwarfed Purdue's spending on MS Contin or
 6 Defendant Janssen's spending on Duragesic:³⁰

Figure 1: Promotional Spending for Three Opioid Analgesics in First 6 Years of Sales



21 98. Prior to Purdue's launch of OxyContin, no drug company had ever promoted such
 22 a pure, high-strength Schedule II narcotic to so wide an audience of general practitioners. Today,
 23 one in every five patients who present themselves to physicians' offices with non-cancer pain
 24

25 ²⁹ *Id.* at 156.

26 ³⁰ *OxyContin Abuse and Diversion and Efforts to Address the Problem*, U.S. Gen. Acct. Off. Rep. to Cong. Requesters at 22 (Dec. 2003), <http://www.gao.gov/new.items/d04110.pdf>.

1 symptoms or pain-related diagnoses (including acute and chronic pain) receives an opioid
 2 prescription.³¹

3 99. Purdue has generated estimated sales of more than \$35 billion from opioids since
 4 1996, while raking in more than \$3 billion in 2015 alone. Remarkably, its opioid sales continued
 5 to climb even after a period of media attention and government inquiries regarding OxyContin
 6 abuse in the early 2000s and a criminal investigation culminating in guilty pleas in 2007. Purdue
 7 proved itself skilled at evading full responsibility and continuing to sell through the controversy.
 8 The company's annual opioid sales of \$3 billion in 2015 represent a four-fold increase from its
 9 2006 sales of \$800 million.

10 100. One might imagine that Richard Sackler's ambitions have been realized. But in
 11 the best tradition of family patriarch Arthur Sackler, Purdue has its eyes on even greater profits.
 12 Under the name of Mundipharma, the Sacklers are looking to new markets for their opioids—
 13 employing the exact same playbook in South America, China, and India as they did in the United
 14 States.

15 101. In May 2017, a dozen members of Congress sent a letter to the World Health
 16 Organization, warning it of the deceptive practices Purdue is unleashing on the rest of the world
 17 through Mundipharma:

18 We write to warn the international community of the deceptive and dangerous
 19 practices of Mundipharma International—an arm of Purdue Pharmaceuticals. The
 20 greed and recklessness of one company and its partners helped spark a public health
 21 crisis in the United States that will take generations to fully repair. We urge the
 22 World Health Organization (WHO) to do everything in its power to avoid allowing
 the same people to begin a worldwide opioid epidemic. Please learn from our
 experience and do not allow Mundipharma to carry on Purdue's deadly legacy on
 a global stage. . . .

23 Internal documents revealed in court proceedings now tell us that since the early
 24 development of OxyContin, Purdue was aware of the high risk of addiction it
 carried. Combined with the misleading and aggressive marketing of the drug by its

25 ³¹ Deborah Dowell, M.D., Tamara M. Haegerich, Ph.D., and Roger Chou, M.D., *CDC Guideline for Prescribing*
 26 *Opioids for Chronic Pain — United States, 2016*, Ctrs. for Disease Control and Prevention (Mar. 18, 2016),
<https://www.cdc.gov/mmwr/volumes/65/rr/rr6501e1.htm> [hereinafter 2016 CDC Guideline].

partner, Abbott Laboratories, Purdue began the opioid crisis that has devastated American communities since the end of the 1990s. Today, Mundipharma is using many of the same deceptive and reckless practices to sell OxyContin abroad. . . .

In response to the growing scrutiny and diminished U.S. sales, the Sacklers have simply moved on. On December 18, the Los Angeles Times published an extremely troubling report detailing how in spite of the scores of lawsuits against Purdue for its role in the U.S. opioid crisis, and tens of thousands of overdose deaths, Mundipharma now aggressively markets OxyContin internationally. In fact, Mundipharma uses many of the same tactics that caused the opioid epidemic to flourish in the U.S., though now in countries with far fewer resources to devote to the fallout.³²

102. Purdue's pivot to untapped markets, after extracting substantial profits from communities like Jefferson County and leaving the County to address the resulting damage, underscores that its actions have been knowing, intentional, and motivated by profits throughout this entire tragic story.

B. The Booming Business of Addiction

1. Other Manufacturing Defendants leapt at the opioid opportunity.

103. Purdue created a market in which the prescription of powerful opioids for a range of common aches and pains was not only acceptable but encouraged—but it was not alone. Defendants Endo, Janssen, Cephalon, and Actavis, each of which already produced and sold prescription opioids, positioned themselves to take advantage of the opportunity Purdue created, developing both branded and generic opioids to compete with OxyContin while misrepresenting the safety and efficacy of their products.

104. Endo, which for decades had sold Percocet and Percodan, both containing relatively low doses of oxycodone, moved quickly to develop a generic version of extended-release oxycodone to compete with OxyContin, receiving tentative FDA approval for its generic version in 2002. As Endo stated in its 2003 Form 10-K, it was the first to file an application with the FDA for bioequivalent versions of the 10, 20, and 40 mg strengths of OxyContin, which

³² Letter from Cong. of the U.S., to Dr. Margaret Chan, Dir.-Gen., World Health Org. (May 3, 2017), <http://katherineclark.house.gov/cache/files/a577bd3c-29ec-4bb9-bdba-1ca71c784113/mundipharma-letter-signatures.pdf>.

1 potentially entitled it to 180 days of generic marketing exclusivity—“a significant advantage.”³³
 2 Purdue responded by suing Endo for patent infringement, litigating its claims through a full trial
 3 and a Federal Circuit appeal—unsuccessfully. As the trial court found, and the appellate court
 4 affirmed, Purdue obtained the oxycodone patents it was fighting to enforce through “inequitable
 5 conduct”—namely, suggesting that its patent applications were supported by clinical data when
 6 in fact they were based on an employee’s “insight and not scientific proof.”³⁴ Endo began selling
 7 its generic extended-release oxycodone in 2005.

8 105. At the same time as Endo was battling Purdue over generic OxyContin—and as
 9 the U.S. was battling increasingly widespread opioid abuse—Endo was working on getting
 10 another branded prescription opioid on the market. In 2002, Endo submitted applications to the
 11 FDA for both immediate-release and extended-release tablets of oxymorphone, branded as
 12 Opana and Opana ER.

13 106. Like oxycodone, oxymorphone is not a new drug; it was first synthesized in
 14 Germany in 1914 and sold in the U.S. by Endo beginning in 1959 under the trade name
 15 Numorphan, in injectable, suppository, and oral tablet forms. But the oral tablets proved highly
 16 susceptible to abuse. Called “blues” after the light blue color of the 10 mg pills, Numorphan
 17 provoked, according to some users, a more euphoric high than heroin, and even had its moment
 18 in the limelight as the focus of the movie *Drugstore Cowboy*. As the National Institute on Drug
 19 Abuse observed in its 1974 report, “Drugs and Addict Lifestyle,” Numorphan was extremely
 20 popular among addicts for its quick and sustained effect.³⁵ Endo withdrew oral Numorphan from
 21 the market in 1979, reportedly for “commercial reasons.”³⁶

22
 23
 24 ³³ *Endo Pharm. Holdings, Inc. Form 10-K*, U.S. Sec. and Exchange Comm’n, at 4 (Mar. 15, 2004),
http://media.corporate-ir.net/media_files/irol/12/123046/reports/10K_123103.pdf.

25 ³⁴ *Purdue Pharma L.P. v. Endo Pharm. Inc.*, 438 F.3d 1123, 1131 (Fed. Cir. 2006).

26 ³⁵ John Fauber and Kristina Fiore, *Abandoned Painkiller Makes a Comeback*, MedPage Today (May 10, 2015),
<https://www.medpagetoday.com/psychiatry/addictions/51448>.

³⁶ *Id.*

107. Two decades later, however, as communities around the U.S. were first sounding the alarm about prescription opioids and Purdue executives were being called to testify before Congress about the risks of OxyContin, Endo essentially reached back into its inventory, dusted off a product it had previously shelved after widespread abuse, and pushed it into the marketplace with a new trade name and a potent extended-release formulation.

108. The clinical trials submitted with Endo's first application for approval of Opana were insufficient to demonstrate efficacy, and some subjects in the trials overdosed and had to be revived with naloxone. Endo then submitted new "enriched enrollment" clinical trials, in which trial subjects who do not respond to the drug are excluded from the trial, and obtained approval. Endo began marketing Opana and Opana ER in 2006.

109. Like Numorphan, Opana ER was highly susceptible to abuse. On June 8, 2017, the FDA sought removal of Opana ER. In its press release, the FDA indicated that "the agency is seeking removal based on its concern that the benefits of the drug may no longer outweigh its risks. This is the first time the agency has taken steps to remove a currently marketed opioid pain medication from sale due to the public health consequences of abuse."³⁷ On July 6, 2017, Endo agreed to withdraw Opana ER from the market.³⁸

110. Janssen, which already marketed the Duragesic (fentanyl) patch, developed a new opioid compound called tapentadol in 2009, marketed as Nucynta for the treatment of moderate to severe pain. Janssen launched the extended-release version, Nucynta ER, for treatment of chronic pain in 2011.

111. Cephalon also manufactures Actiq, a fentanyl lozenge, and Fentora, a fentanyl tablet. As noted above, fentanyl is an extremely powerful synthetic opioid. According to the DEA, as little as two milligrams is a lethal dosage for most people. Actiq has been approved by

³⁷ Press Release, U.S. Food & Drug Administration, *FDA requests removal of Opana ER for risks related to abuse* (June 8, 2017), <https://www.fda.gov/NewsEvents/Newsroom/PressAnnouncements/ucm562401.htm>.

³⁸ *Endo pulls opioid as U.S. seeks to tackle abuse epidemic*, Reuters (July 6, 2017, 9:59am), <https://www.reuters.com/article/us-endo-intl-opana-idUSKBN19R2II>.

1 the FDA only for the “management of breakthrough cancer pain in patients 16 years and older
 2 with malignancies who are already receiving and who are tolerant to around-the-clock opioid
 3 therapy for the underlying persistent cancer pain.”³⁹ Fentora has been approved by the FDA only
 4 for the “management of breakthrough pain in cancer patients 18 years of age and older who are
 5 already receiving and who are tolerant to around-the-clock opioid therapy for their underlying
 6 persistent cancer pain.”⁴⁰

7 112. In 2008, Cephalon pled guilty to a criminal violation of the Federal Food, Drug
 8 and Cosmetic Act for its misleading promotion of Actiq and two other drugs and agreed to pay
 9 \$425 million.

10 113. Actavis acquired the rights to Kadian, extended-release morphine, in 2008, and
 11 began marketing Kadian in 2009. Actavis’s opioid products also include Norco, a brand-name
 12 hydrocodone and acetaminophen pill, first approved in 1997. But Actavis, primarily a generic
 13 drugmaker, pursued opioid profits through generics, selling generic versions of OxyContin,
 14 Opana, and Duragesic. In 2013, it settled a patent lawsuit with Purdue over its generic version of
 15 “abuse-deterrent” OxyContin, striking a deal that would allow it to market its abuse-deterrent
 16 oxycodone formulation beginning in 2014. Actavis anticipated over \$100 million in gross profit
 17 from generic OxyContin sales in 2014 and 2015.

18 114. Mallinckrodt’s generic oxycodone achieved enough market saturation to have its
 19 own street name, “M’s,” based on its imprint on the pills. As noted above, Mallinckrodt was the
 20 subject of a federal investigation based on diversion of its oxycodone in Florida, where 500
 21 million of its pills were shipped between 2008 and 2012. Federal prosecutors alleged that 43,991
 22 orders from distributors and retailers were excessive enough be considered suspicious and should
 23 have been reported to the DEA.

24
 25 ³⁹ *Prescribing Information, ACTIQ®*, U.S. Food & Drug Admin.,
https://www.accessdata.fda.gov/drugsatfda_docs/label/2009/020747s030lbl.pdf (last visited Aug. 13, 2018).

26 ⁴⁰ *Prescribing Information, FENTORA®*, U.S. Food & Drug Admin.,
https://www.accessdata.fda.gov/drugsatfda_docs/label/2012/021947s015lbl.pdf (last visited Aug. 13, 2018).

115. Mallinckrodt also pursued a share of the branded opioid market. In 2007, Mallinckrodt launched Magnacet, an oxycodone and acetaminophen combination pill. In 2008, Mallinckrodt (then Covidien) launched TussiCaps, a hydrocodone and chlorpheniramine capsule, marketed as a cough suppressant. And in 2009, Mallinckrodt acquired the U.S. rights to Exalgo, a potent extended-release hydromorphone tablet, and began marketing it in 2012. Mallinckrodt further expanded its branded opioid portfolio in 2012 by purchasing Roxicodone from Xanodyne Pharmaceuticals. In addition, Mallinckrodt developed Xartemis XR, an extended-release combination of oxycodone and acetaminophen, which the FDA approved in March 2014. In anticipation of Xartemis XR's approval, Mallinckrodt hired approximately 200 sales representatives to promote it, and CEO Mark Trudeau said the drug could generate "hundreds of millions in revenue."⁴¹

116. All told, the Manufacturing Defendants have reaped enormous profits from the addiction crisis they spawned. For example, Opana ER alone generated more than \$1 billion in revenue for Endo in 2010 and again in 2013. Janssen earned more than \$1 billion in sales of Duragesic in 2009, and Nucynta and Nucynta ER accounted for \$172 million in sales in 2014.

2. Distributor Defendants knowingly supplied dangerous quantities of opioids while advocating for limited oversight and enforcement.

117. The Distributor Defendants track and keep a variety of information about the pharmacies and other entities to which they sell pharmaceuticals. For example, the Distributor Defendants use "know your customer" questionnaires that track the number and types of pills their customers sell, absolute and relative amounts of controlled substances they sell, whether the customer purchases from other distributors, and types of medical providers in the areas, among other information.

⁴¹ Samantha Liss, *Mallinckrodt banks on new painkillers for sales*, St. Louis Bus. Journal (Dec. 30, 2013), <http://argencapital.com/mallinckrodt-banks-on-new-painkillers-for-sales/>.

1 118. These questionnaires and other sources of information available to the Distributor
2 Defendants provide ample data to put the Distributor Defendants on notice of suspicious orders,
3 pharmacies, and doctors.

4 119. Nevertheless, the Distributor Defendants refused or failed to identify, investigate,
5 or report suspicious orders of opioids to the DEA. Even when the Distributor Defendants had
6 actual knowledge that they were distributing opioids to drug diversion rings, they refused or
7 failed to report these sales to the DEA.

8 120. By not reporting suspicious opioid orders or known diversions of prescription
9 opioids, not only were the Defendants able to continue to sell opioids to questionable customers,
10 Defendants ensured that the DEA had no basis for decreasing or refusing to increase production
11 quotas for prescription opioids.

12 121. The Distributor Defendants collaborated with each other and with the
13 Manufacturing Defendants to maintain distribution of excessive amounts of opioids. One
14 example of this collaboration came to light through Defendants' work in support of legislation
15 called the Ensuring Patient Access and Effective Drug Enforcement (EPAEDE) Act, which was
16 signed into law in 2016 and limited the DEA's ability to stop the flow of opioids. Prior to this
17 law, the DEA could use an "immediate suspension order" to halt suspicious shipments of pills
18 that posed an "imminent" threat to the public. The EPAEDE Act changed the required showing
19 to an "immediate" threat—an impossible standard given the fact that the drugs may sit on a shelf
20 for a few days after shipment. The law effectively neutralized the DEA's ability to bring
21 enforcement actions against distributors.

22 122. The legislation was drafted by a former DEA lawyer, D. Linden Barber, who is
23 now a senior vice president at Defendant Cardinal Health. Prior to leaving the DEA, Barber had
24 worked with Joseph Rannazzisi, then the chief of the DEA's Office of Diversion Control, to plan
25 the DEA's fight against the diversion of prescription drugs. So when Barber began working for
26 Cardinal Health, he knew just how to neutralize the effectiveness of the DEA's enforcement

1 actions. Barber and other promoters of the EPAEDE Act portrayed the legislation as maintaining
 2 patient access to medication critical for pain relief. In a 2014 hearing on the bill, Barber testified
 3 about the “unintended consequences in the supply chain” of the DEA’s enforcement actions. But
 4 by that time, communities across the United States, including Plaintiff Jefferson County, were
 5 grappling with the “unintended consequences” of Defendants’ reckless promotion and
 6 distribution of narcotics.

7 123. Despite egregious examples of drug diversion from around the country, the
 8 promoters of the EPAEDE Act were successful in characterizing the bill as supporting patients’
 9 rights. One of the groups supporting this legislation was the Alliance for Patient Access, a “front
 10 group” as discussed further below, which purports to advocate for patients’ rights to have access
 11 to medicines, and whose 2017 list of “associate members and financial supporters” included
 12 Defendants Purdue, Endo, Johnson & Johnson, Actavis, Mallinckrodt, and Cephalon. In a 2013
 13 “white paper” titled “Prescription Pain Medication: Preserving Patient Access While Curbing
 14 Abuse,” the Alliance for Patient Access asserted multiple “unintended consequences” of
 15 regulating pain medication, including a decline in prescriptions as physicians feel burdened by
 16 regulations and stigmatized.⁴²

17 124. The Distributor Defendants are also part of the activities of the Alliance for
 18 Patient Access, although their involvement is hidden. One example of their involvement was
 19 revealed by the metadata of an electronic document: the letter from the Alliance for Patient
 20 Access in support of the EPAEDE Act. That document was created by Kristen Freitas, a
 21 registered lobbyist and the vice president for federal government affairs of the Healthcare
 22 Distributors Alliance (HDA)—the trade group that represents Defendants McKesson, Cardinal
 23 Health, and AmerisourceBergen.

25 ⁴² *Prescription Pain Medication: Preserving Patient Access While Curbing Abuse*, Inst. for Patient Access (Oct.
 26 2013), http://1yh21u3cjptv3xjder1dco9mx5s.wpengine.netdna-cdn.com/wp-content/uploads/2013/12/PT_White-Paper_Finala.pdf.

125. Upon information and belief, the collaboration on the EPAEDE Act is just one example of how the Manufacturing Defendants and the Distributor Defendants, through third-party “front groups” like the Alliance for Patient Access and trade organizations like HDA, worked together behind the scenes to ensure that the flow of dangerous narcotics into communities across the country would not be restricted, and Defendants collaborated in other ways that remain hidden from public view.

126. The Distributor Defendants have been the subject of numerous enforcement actions by the DEA. In 2008, for example, McKesson was fined \$13.3 million and agreed to strengthen its controls by implementing a three-tiered system that would flag buyers who exceeded monthly thresholds for opioids. As the opioid crisis deepened, the DEA’s Office of Diversion Control, led by Rannazzisi, stepped up enforcement, filing 52 immediate suspension orders against suppliers and pill mills in 2010 alone. Defendant Cardinal Health was fined \$34 million by the DEA in 2013 for failing to report suspicious orders.

127. The Distributor Defendants were not simply passive transporters of opioids. They intentionally failed to report suspicious orders and actively pushed back against efforts to enforce the law and restrict the flow of opioids into communities like Jefferson County.

3. Pill mills and overprescribing doctors also placed their financial interests ahead of their patients’ interests.

128. Prescription opioid manufacturers and distributors were not the only ones to recognize an economic opportunity. Around the country, including in Jefferson County, certain doctors or pain clinics ended up doing brisk business dispensing opioid prescriptions. As Dr. Andrew Kolodny, cofounder of Physicians for Responsible Opioid Prescribing, observed, this business model meant doctors would “have a practice of patients who’ll never miss an appointment and who pay in cash.”⁴³

⁴³ Sam Quinones, *Dreamland: The True Tale of America’s Opiate Epidemic* 314 (Bloomsbury Press 2015).

129. Moreover, the Manufacturing Defendants' sales incentives rewarded sales representatives who happened to have pill mills within their territories, enticing those representatives to look the other way even when their in-person visits to such clinics should have raised numerous red flags. In one example, a pain clinic in South Carolina was diverting massive quantities of OxyContin. People traveled to the clinic from towns as far as 100 miles away to get prescriptions. Eventually, the DEA's diversion unit raided the clinic, and prosecutors filed criminal charges against the doctors. But Purdue's sales representative for that territory, Eric Wilson, continued to promote OxyContin sales at the clinic. He reportedly told another local physician that this clinic accounted for 40% of the OxyContin sales in his territory. At that time, Wilson was Purdue's top-ranked sales representative.⁴⁴ In response to news stories about this clinic, Purdue issued a statement, declaring that "if a doctor is intent on prescribing our medication inappropriately, such activity would continue regardless of whether we contacted the doctor or not."⁴⁵

130. Another pill mill, this one in Los Angeles, supplied OxyContin to a drug dealer in Everett, Washington. Purdue was alerted to the existence of this pill mill by one of its regional sales managers, who in 2009 reported to her supervisors that when she visited the clinic with her sales representative, "it was packed with a line out the door, with people who looked like gang members," and that she felt "very certain that this an organized drug ring[.]" She wrote, "This is clearly diversion. Shouldn't the DEA be contacted about this?" But her supervisor at Purdue responded that while they were "considering all angles," it was "really up to [the wholesaler] to make the report." This clinic was the source of 1.1 million pills trafficked to Everett, which is a city of around 100,000 people. Purdue waited until after the clinic was shut down in 2010 to inform the authorities.⁴⁶ Similarly, Purdue received repeated reports in 2008 from a sales

⁴⁴ Meier, *supra* note 15, at 298-300.

⁴⁵ *Id.*

⁴⁶ Harriet Ryan, Scott Glover, and Lisa Girion, *How black-market OxyContin spurred a town's descent into crime, addiction and heartbreak*, Los Angeles Times (July 10, 2016), <http://www.latimes.com/projects/la-me-oxycontin-everett/>; Harriet Ryan, Lisa Girion, and Scott Glover, *More than 1 million OxyContin pills ended up in the hands*

1 representative who visited a family practice doctor in Bothell, Washington; the sales
 2 representative informed Purdue that many of this doctor's patients were men in their twenties
 3 who did not appear to be in pain, who sported diamond studs and \$350 sneakers, and who always
 4 paid for their 80 mg OxyContin prescriptions in cash. Despite being repeatedly alerted to the
 5 doctor's conduct, Purdue did not take any action to report it until three years later.

6 131. Whenever examples of opioid diversion and abuse have drawn media attention,
 7 the Manufacturing Defendants have consistently blamed "bad actors." For example, in 2001,
 8 during a Congressional hearing, Purdue's attorney Howard Udell answered pointed questions
 9 about how it was that Purdue could utilize IMS Health data to assess their marketing efforts but
 10 not notice a particularly egregious pill mill in Pennsylvania run by a doctor named Richard
 11 Paolino. Udell asserted that Purdue was "fooled" by the "bad actor" doctor: "The picture that is
 12 painted in the newspaper [of Dr. Paolino] is of a horrible, bad actor, someone who preyed upon
 13 this community, who caused untold suffering. And he fooled us all. He fooled law enforcement.
 14 He fooled the DEA. He fooled local law enforcement. He fooled us."⁴⁷

15 132. But given the closeness with which all Defendants monitored prescribing patterns,
 16 including through IMS Health data, it is highly improbable that they were "fooled." In fact, a
 17 local pharmacist had noticed the volume of prescriptions coming from Paolino's clinic and
 18 alerted authorities. Purdue had the prescribing data from the clinic and alerted no one. Rather, it
 19 appears Purdue and other Defendants used the IMS Health data to target pill mills and sell more
 20 pills. Indeed, a Purdue executive referred to Purdue's tracking system and database as a "gold
 21 mine" and acknowledged that Purdue could identify highly suspicious volumes of prescriptions.

22 133. Sales representatives making in-person visits to such clinics were likewise not
 23 fooled. But as pill mills were lucrative for the manufacturers and individual sales representatives
 24 alike, Defendants and their employees turned a collective blind eye, allowing certain clinics to

25 *of criminals and addicts. What the drugmaker knew*, Los Angeles Times (July 10, 2016),
 26 <http://www.latimes.com/projects/la-me-oxycontin-part2/>.

⁴⁷ Meier, *supra* note 15, at 179.

1 dispense staggering quantities of potent opioids and feigning surprise when the most egregious
2 examples eventually made the nightly news.

3 **4. Widespread prescription opioid use broadened the market for heroin and**
4 **fentanyl.**

5 134. Defendants' scheme achieved a dramatic expansion of the U.S. market for
6 opioids, prescription and non-prescription alike. Heroin and fentanyl use has surged—a
7 foreseeable consequence of Defendants' successful promotion of opioid use coupled with the
8 sheer potency of their products.

9 135. In his book *Dreamland: The True Tale of America's Opiate Epidemic*, journalist
10 Sam Quinones summarized the easy entrance of black tar heroin in a market primed by
11 prescription opioids:

12 His black tar, once it came to an area where OxyContin had already tenderized the
13 terrain, sold not to tapped-out junkies but to younger kids, many from the suburbs,
14 most of whom had money and all of whom were white. Their transition from Oxy
15 to heroin, he saw, was a natural and easy one. Oxy addicts began by sucking on and
16 dissolving the pills' timed-release coating. They were left with 40 or 80 mg of pure
17 oxycodone. At first, addicts crushed the pills and snorted the powder. As their
18 tolerance built, they used more. To get a bigger bang from the pill, they liquefied it
and injected it. But their tolerance never stopped climbing. OxyContin sold on the
street for a dollar a milligram and addicts very quickly were using well over 100
mg a day. As they reached their financial limits, many switched to heroin, since
they were already shooting up Oxy and had lost any fear of the needle.⁴⁸

19 136. In a study examining the relationship between the abuse of prescription opioids
20 and heroin, researchers found that 75% of those who began their opioid abuse in the 2000s
21 reported that their first opioid was a prescription drug.⁴⁹ As the graph below illustrates,
22 prescription opioids replaced heroin as the first opioid of abuse beginning in the 1990s.
23

24
25 ⁴⁸ Quinones, *supra* note 43, at 165-66.

26 ⁴⁹ Theodore J. Cicero, PhD, Matthew S. Ellis, MPE, Hilary L. Surratt, PhD, *The Changing Face of Heroin Use in the United States: A Retrospective Analysis of the Past 50 Years*, 71(7) JAMA Psychiatry 821-826 (2014), <https://jamanetwork.com/journals/jamapsychiatry/fullarticle/1874575>.



From: **The Changing Face of Heroin Use in the United States: A Retrospective Analysis of the Past 50 Years**

JAMA Psychiatry. 2014;71(7):821-826. doi:10.1001/jamapsychiatry.2014.366

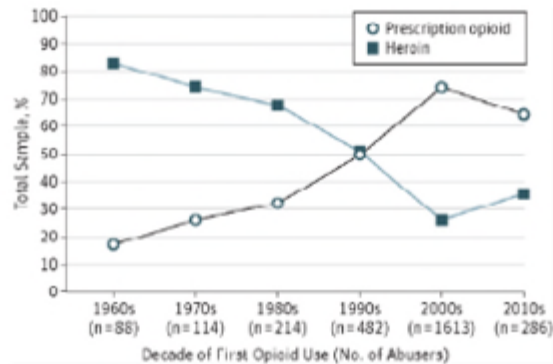


Figure Legend:

Percentage of the Total Heroin-Dependent Sample That Used Heroin or a Prescription Opioid as Their First Opioid of Abuse Data are plotted as a function of the decade in which respondents initiated their opioid abuse.

137. The researchers also found that nearly half of the respondents who indicated that their primary drug was heroin actually preferred prescription opioids, because the prescription drugs were legal, and perceived as “safer and cleaner.” But, heroin’s lower price point is a distinct advantage. While an 80 mg OxyContin might cost \$80 on the street, the same high can be had from \$20 worth of heroin.

138. As noted above, there is little difference between the chemical structures of heroin and prescription opioids. Between 2005 and 2009, Mexican heroin production increased by over 600%. And between 2010 and 2014, the amount of heroin seized at the U.S.-Mexico border more than doubled.

1 139. From 2002 to 2016, fatal overdoses related to heroin in the U.S. increased by
2 **533%**—from 2,089 deaths in 2002 to 13,219 deaths in 2016.⁵⁰

3 140. Along with heroin use, fentanyl use is on the rise, as a result of America's
4 expanded appetite for opioids. But fentanyl, as noted above, is fifty times more potent than
5 heroin, and overdosing is all too easy. Fentanyl is expected to cause over 20,000 overdoses in
6 2017.⁵¹

7 141. As Dr. Caleb Banta-Green, senior research scientist at the University of
8 Washington's Alcohol and Drug Abuse Institute, told The Seattle Times in August 2017, "The
9 bottom line is opioid addiction is the overall driver of deaths. People will use whatever opioid
10 they can get. It's just that which one they're buying is changing a bit."⁵²

11 142. In addition to the expanded market for opioids of all kinds, the opioid epidemic
12 has contributed to a resurgence in methamphetamine use, as some opioid users turn to the
13 stimulant to counter the effects of opioids.⁵³ As explained in a recent article regarding the
14 connection between opioids and methamphetamine, "[f]or addicts, the drugs pair: Heroin is a
15 downer and methamphetamine is an upper."⁵⁴

16 **C. The Manufacturing Defendants Promoted Prescription Opioids Through Several**
17 **Channels.**

18 143. Despite knowing the devastating consequences of widespread opioid use, the
19 Manufacturing Defendants engaged in a sophisticated and multi-pronged promotional campaign

20
21 ⁵⁰ Niall McCarthy, *U.S. Heroin Deaths Have Increased 533% Since 2002*, Forbes (Sept. 11, 2017, 8:26am),
22 <https://www.forbes.com/sites/niallmccarthy/2017/09/11/u-s-heroin-deaths-have-increased-533-since-2002-infographic/#13ab9a531abc>.

⁵¹ *Id.*

23 ⁵² *Opioids: The Leading Cause of Drug Deaths in Seattle Area*, U. of Wash. Sch. of Pub. Health (Aug. 25, 2017),
24 http://sph.washington.edu/news/article.asp?content_ID=8595.

25 ⁵³ See, e.g., *Opioids and methamphetamine: a tale of two crises*, 391(10122) The Lancet 713 (Feb. 24, 2018),
26 [https://www.thelancet.com/journals/lancet/article/PIIS0140-6736\(18\)30319-2/fulltext](https://www.thelancet.com/journals/lancet/article/PIIS0140-6736(18)30319-2/fulltext); Brenda Goodman, MA,
Experts Warn of Emerging 'Stimulant Epidemic', WebMD (Apr. 3, 2018), <https://www.webmd.com/mental-health/addiction/news/20180403/experts-warn-of-emerging-stimulant-epidemic>.

⁵⁴ Michelle Theriault Boots, *The silent fallout of the opioid epidemic? Meth.*, Anchorage Daily News (Mar. 29, 2018), <https://www.adn.com/alaska-news/2018/03/19/the-silent-fallout-of-the-opioid-epidemic-meth/#>.

designed to achieve just that. By implementing the strategies pioneered by Arthur Sackler, these Defendants were able to achieve the fundamental shift in the perception of opioids that was key to making them blockbuster drugs.

144. The Manufacturing Defendants disseminated their deceptive statements about opioids through several channels.⁵⁵ First, these Defendants aggressively and persistently pushed opioids through sales representatives. Second, these Defendants funded third-party organizations that appeared to be neutral, but which served as additional marketing departments for drug companies. Third, these Defendants utilized prominent physicians as paid spokespeople—“Key Opinion Leaders”—to take advantage of doctors’ respect for and reliance on the recommendations of their peers. Finally, these Defendants also used print and online advertising, including unbranded advertising, which is not reviewed by the FDA.

145. The Manufacturing Defendants spent substantial sums and resources in making these communications. For example, Purdue spent more than \$200 million marketing OxyContin in 2001 alone.⁵⁶

1. The Manufacturing Defendants aggressively deployed sales representatives to push their products.

146. The Manufacturing Defendants communicated to prescribers directly in the form of in-person visits and communications from sales representatives.

147. The Manufacturing Defendants’ tactics through their sales representatives—also known as “detailers”—were particularly aggressive. In 2014, Manufacturing Defendants collectively spent well over \$100 million on detailing branded opioids to doctors.

⁵⁵ The specific misrepresentations and omissions are discussed below in Section D.

⁵⁶ *Oxycontin: Balancing Risks and Benefits: Hearing of the S. Comm. on Health, Education, Labor and Pensions*, 107th Cong. 2 (Feb. 12, 2002) (testimony of Paul Goldenheim, Vice President for Research, Purdue Pharma), <https://www.gpo.gov/fdsys/pkg/CHRG-107shrg77770/html/CHRG-107shrg77770.htm>.

148. Each sales representative has a specific sales territory and is responsible for developing a list of about 105 to 140 physicians to call on who already prescribe opioids or who are candidates for prescribing opioids.

149. When Purdue launched OxyContin in 1996, its 300-plus sales force had a total physician call list of approximately 33,400 to 44,500. By 2000, nearly 700 representatives had a total call list of approximately 70,500 to 94,000 physicians. Each sales representative was expected to make about thirty-five physician visits per week and typically called on each physician every three to four weeks, while each hospital sales representative was expected to make about fifty physician visits per week and call on each facility every four weeks.⁵⁷

150. One of Purdue's early training memos compared doctor visits to "firing at a target," declaring that "[a]s you prepare to fire your 'message,' you need to know where to aim and what you want to hit!"⁵⁸ According to the memo, the target is physician resistance based on concern about addiction: "The physician wants pain relief for these patients without addicting them to an opioid."⁵⁹

151. Former sales representative Steven May, who worked for Purdue from 1999 to 2005, explained to a journalist that the most common objection he heard about prescribing OxyContin was that "it's just too addictive."⁶⁰ In order to overcome that objection and hit their "target," May and other sales representatives were taught to say, "The delivery system is believed to reduce the abuse liability of the drug."⁶¹ May repeated that line to doctors even

⁵⁷ *OxyContin Abuse and Diversion and Efforts to Address the Problem*, *supra* note 30, at 20.

⁵⁸ Meier, *supra* note 15, at 102.

⁵⁹ *Id.*

⁶⁰ David Remnick, *How OxyContin Was Sold to the Masses* (Steven May interview with Patrick Radden Keefe), *New Yorker* (Oct. 27, 2017), <https://www.newyorker.com/podcast/the-new-yorker-radio-hour/how-oxycontin-was-sold-to-the-masses>.

⁶¹ Patrick Radden Keefe, *The Family That Built an Empire of Pain*, *New Yorker* (Oct. 30, 2017), <https://www.newyorker.com/magazine/2017/10/30/the-family-that-built-an-empire-of-pain>; see also Meier, *supra* note 15, at 102 ("Delayed absorption, as provided by OxyContin tablets, is believed to reduce the abuse liability of the drug.").

1 though he “found out pretty fast that it wasn’t true.”⁶² He and his coworkers learned quickly that
 2 people were figuring out how to remove the time-releasing coating, but they continued making
 3 this misrepresentation until Purdue was forced to remove it from the drug’s label.

4 152. Purdue trained its sales representatives to misrepresent the addiction risk in other
 5 ways. May explained that he and his coworkers were trained to “refocus” doctors on “legitimate”
 6 pain patients, and to represent that “legitimate” patients would not become addicted. In addition,
 7 they were trained to say that the 12-hour dosing made the extended-release opioids less “habit-
 8 forming” than painkillers that need to be taken every four hours. Similarly, former Purdue sales
 9 manager William Gergely told a Florida state investigator in 2002 that sales representatives were
 10 instructed to say that OxyContin was “virtually non-addicting” and “non-habit-forming.”⁶³

11 153. As Shelby Sherman, a Purdue sales representative from 1974 to 1998, told a
 12 reporter regarding OxyContin promotion, “It was sell, sell, sell. We were directed to lie. Why
 13 mince words about it?”⁶⁴

14 154. The Manufacturing Defendants utilized lucrative bonus systems to encourage
 15 their sales representatives to stick to the script and increase opioid sales in their territories.
 16 Purdue paid \$40 million in sales incentive bonuses to its sales representatives in 2001 alone, with
 17 annual bonuses ranging from \$15,000 to nearly \$240,000.⁶⁵ The training memo described above,
 18 in keeping with a Wizard of Oz theme, reminded sales representatives: “A pot of gold awaits you
 19 ‘Over the Rainbow’!”⁶⁶

20 155. As noted above, these Defendants have also spent substantial sums to purchase,
 21 manipulate, and analyze prescription data available from IMS Health, which allows them to track

22 ⁶² Keefe, *supra* note 61.

23 ⁶³ Fred Schulte and Nancy McVicar, *Oxycontin Was Touted As Virtually Nonaddictive, Newly Released State*
 24 *Records Show*, Sun Sentinel (Mar. 6, 2003), http://articles.sun-sentinel.com/2003-03-06/news/0303051301_1_purdue-pharma-oxycontin-william-gergely.

25 ⁶⁴ Glazek, *supra* note 26.

26 ⁶⁵ Art Van Zee, M.D., *The Promotion and Marketing of OxyContin: Commercial Triumph, Public Health Tragedy*,
 99(2) Am J Public Health 221-27 (Feb. 2009), <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC2622774/>.

⁶⁶ Meier, *supra* note 15, at 103.

1 initial prescribing and refill practices by individual doctors, and in turn to customize their
2 communications with each doctor. The Manufacturing Defendants' use of this marketing data
3 was a cornerstone of their marketing plan,⁶⁷ and continues to this day.

4 156. The Manufacturing Defendants also aggressively pursued family doctors and
5 primary care physicians perceived to be susceptible to their marketing campaigns. The
6 Manufacturing Defendants knew that these doctors relied on information provided by
7 pharmaceutical companies when prescribing opioids, and that, as general practice doctors seeing
8 a high volume of patients on a daily basis, they would be less likely to scrutinize the companies'
9 claims.

10 157. Furthermore, the Manufacturing Defendants knew or should have known the
11 doctors they targeted were often poorly equipped to treat or manage pain comprehensively, as
12 they often had limited resources or time to address behavioral or cognitive aspects of pain
13 treatment or to conduct the necessary research themselves to determine whether opioids were as
14 beneficial as these Defendants claimed. In fact, the majority of doctors and dentists who
15 prescribe opioids are not pain specialists. For example, a 2014 study conducted by pharmacy
16 benefit manager Express Scripts reviewing narcotic prescription data from 2011 to 2012
17 concluded that of the more than 500,000 prescribers of opioids during that time period, *only* 385
18 were identified as pain specialists.⁶⁸

19 158. When the Manufacturing Defendants presented these doctors with sophisticated
20 marketing material and apparently scientific articles that touted opioids' ability to easily and
21 safely treat pain, many of these doctors began to view opioids as an efficient and effective way to
22 treat their patients.

23 159. In addition, sales representatives aggressively pushed doctors to prescribe
24 stronger doses of opioids. For example, one Purdue sales representative in Florida wrote about
25

26 ⁶⁷ Van Zee, *The Promotion and Marketing of OxyContin*, *supra* note 65.

⁶⁸ *A Nation in Pain*, Express Scripts (Dec. 9, 2014), <http://lab.express-scripts.com/lab/publications/a-nation-in-pain>.

1 working for a particularly driven regional manager named Chris Sposato and described how
2 Sposato would drill the sales team on their upselling tactics:

3 It went something like this. "Doctor, what is the highest dose of OxyContin you
4 have ever prescribed?" "20mg Q12h." "Doctor, if the patient tells you their pain
5 score is still high you can increase the dose 100% to 40mg Q12h, will you do that?"
6 "Okay." "Doctor, what if that patient then came back and said their pain score was
7 still high, did you know that you could increase the OxyContin dose to 80mg Q12h,
8 would you do that?" "I don't know, maybe." "Doctor, but you do agree that you
9 would at least Rx the 40mg dose, right?" "Yes."

10 The next week the rep would see that same doctor and go through the same
11 discussion with the goal of selling higher and higher doses of OxyContin. Miami
12 District reps have told me that on work sessions with [Sposato] they would sit in
13 the car and role play for as long as it took until [Sposato] was convinced the rep
14 was delivering the message with perfection.

15 160. The Manufacturing Defendants used not only incentives but competitive pressure
16 to push sales representatives into increasingly aggressive promotion. One Purdue sales
17 representative recalled the following scene: "I remember sitting at a round table with others from
18 my district in a regional meeting while everyone would stand up and state the highest dose that
19 they had suckered a doctor to prescribe. The entire region!!"

20 161. Sales representatives knew that the prescription opioids they were promoting were
21 dangerous. For example, May had only been at Purdue for two months when he found out that a
22 doctor he was calling on had just lost a family member to an OxyContin overdose.⁶⁹ And as
23 another sales representative wrote on a public forum:

24 Actions have consequences - so some patient gets Rx'd the 80mg OxyContin when
25 they probably could have done okay on the 20mg (but their doctor got "sold" on
26 the 80mg) and their teen son/daughter/child's teen friend finds the pill bottle and
takes out a few 80's... next they're at a pill party with other teens and some kid
picks out a green pill from the bowl... they go to sleep and don't wake up (because
they don't understand respiratory depression) Stupid decision for a teen to
make...yes... but do they really deserve to die?

⁶⁹ Remnick, *supra* note 60.

1 162. The Marketing Defendants rewarded their sales representatives with bonuses
 2 when doctors whom they had detailed wrote prescriptions for their company's drug. Because of
 3 this incentive system, sales representatives stood to gain significant bonuses if they had a pill
 4 mill in their sales region.⁷⁰ Sales representatives could be sure that doctors and nurses at pill
 5 mills would be particularly receptive to their messages and incentives, and receive "credit" for
 6 the many prescriptions these pill mills wrote.

7 163. As a result, sales representatives continued to promote opioids even at known pill
 8 mills, including in Washington State, such as Seattle Pain Clinic locations directed by Dr. Frank
 9 Li—who eventually had his medical license suspended for improperly prescribing opioids.
 10 During detailers' frequent visits to Dr. Li, they often noted circumstances that should have led
 11 them to discontinue sales calls and report Dr. Li and his staff to the appropriate authorities.
 12 Instead, they continued to target him for detailing visits that incited him to prescribe even more
 13 opioids, with disastrous consequences for public health.

14 164. In addition, detailers told providers at Dr. Li's clinic that the Washington State
 15 opioid prescription guidelines were wrong and overly conservative, including those related to
 16 calculating the relative strength of different brands of opioids. These detailers often urged
 17 Dr. Li's staff to give patients more opioids, and particular brands of opioids, even when this was
 18 incorrect or conflicted with Washington State guidelines or other medical information.

19 165. Purdue's sales call notes also repeatedly reference how busy Dr. Li and his staff
 20 were—which, combined with the exceptionally high number of opioid prescriptions written by
 21 Dr. Li, should have been another red flag that OxyContin and other opioids were likely being
 22 abused.

23 166. The Manufacturing Defendants' sales representatives also provided health care
 24 providers with pamphlets, visual aids, and other marketing materials designed to increase the rate

25 ⁷⁰ Indeed, Defendants often helped their sales representatives find and target such pill mills. As recently as 2016,
 26 Purdue commissioned a marketing study to help target Washington prescribers and spread its deceptive message
 regarding opioids, and on information and belief, utilized its sales representatives to carry out these strategies.

1 of opioids prescribed to patients. These sales representatives knew the doctors they visited relied
 2 on the information they provided, and that the doctors had minimal time or resources to
 3 investigate the materials' veracity independently.

4 167. The Manufacturing Defendants applied this combination of intense competitive
 5 pressure and lucrative financial incentives because they knew that sales representatives, with
 6 their frequent in-person visits with prescribers, were incredibly effective. In fact, manufacturers'
 7 internal documents reveal that they considered sales representatives their "most valuable
 8 resource."

9 **2. The Manufacturing Defendants bankrolled seemingly independent "front**
 10 **groups" to promote opioid use and fight restrictions on opioids.**

11 168. The Manufacturing Defendants funded, controlled, and operated third-party
 12 organizations that communicated to doctors, patients, and the public the benefits of opioids to
 13 treat chronic pain. These organizations—also known as "front groups"—appeared independent
 14 and unbiased. But in fact, they were but additional paid mouthpieces for the drug manufacturers.
 15 These front groups published prescribing guidelines and other materials that promoted opioid
 16 treatment as a way to address patients' chronic pain. The front groups targeted doctors, patients,
 17 and lawmakers, all in coordinated efforts to promote opioid prescriptions.

18 169. The Manufacturing Defendants spent significant financial resources contributing
 19 to and working with these various front groups to increase the number of opioid prescriptions
 20 written.

21 170. The most prominent front group utilized by the Manufacturing Defendants was
 22 the **American Pain Foundation** (APF), which received more than \$10 million from opioid drug
 23 manufacturers, including Defendants, from 2007 through 2012. For example, Purdue contributed
 24 \$1.7 million and Endo also contributed substantial sums to the APF.⁷¹

25
 26 ⁷¹Charles Ornstein and Tracy Weber, *The Champion of Painkillers*, ProPublica (Dec. 23, 2011, 9:15am),
<https://www.propublica.org/article/the-champion-of-painkillers>.

171. Throughout its existence, APF's operating budget was almost entirely comprised of contributions from prescription opioid manufacturers. For instance, nearly 90% of APF's \$5 million annual budget in 2010 came from "donations" from some of the Manufacturing Defendants, and by 2011, APF was entirely dependent on grants from drug manufacturers, including from Purdue and Endo. Not only did Defendants control APF's purse strings, APF's board of directors was comprised of doctors who were on Defendants' payrolls, either as consultants or speakers at medical events.⁷²

172. Although holding itself out as an independent advocacy group promoting patient well-being, APF consistently lobbied against federal and state proposals to limit opioid use.

173. Another prominent front group was the **American Academy of Pain Medicine** (AAPM), which has received over \$2.2 million in funding since 2009 from opioid drug manufacturers, including Defendants. Like APF, AAPM presented itself as an independent and non-biased advocacy group representing physicians practicing in the field of pain medicine, but in fact was just another mouthpiece the Manufacturing Defendants used to push opioids on doctors and patients.⁷³

174. Both the APF and the AAPM published treatment guidelines and sponsored and hosted medical education programs that touted the benefits of opioids to treat chronic pain while minimizing and trivializing their risks. The treatment guidelines the front groups published—many of which are discussed in detail below—were particularly important to Defendants in ensuring widespread acceptance for opioid therapy to treat chronic pain. Defendants realized, just as the CDC has, that such treatment guidelines can "change prescribing practices," because they appear to be unbiased sources of evidence-based information, even when they are in reality marketing materials.

⁷² *Id.*

⁷³ Tracy Weber and Charles Ornstein, *Two Leaders in Pain Treatment Have Long Ties to Drug Industry*, ProPublica (Dec. 23, 2011, 9:14am), <https://www.propublica.org/article/two-leaders-in-pain-treatment-have-long-ties-to-drug-industry>.

175. For instance, the AAPM, in conjunction with the **American Pain Society** (APS), issued comprehensive guidelines in 2009 titled “Guideline for the Use of Chronic Opioid Therapy in Chronic Noncancer Pain – Evidence Review” (“2009 Guidelines”). The 2009 Guidelines promoted opioids as “safe and effective” for treating chronic pain, despite acknowledging limited evidence to support this statement. Unsurprisingly, the Manufacturing Defendants have widely referenced and promoted these guidelines, issued by front groups these Defendants funded and controlled. These 2009 Guidelines are still available online today.⁷⁴

176. The **Alliance for Patient Access** (APA), discussed above, was established in 2006, along with the firm that runs it, Woodberry Associates LLC. The APA describes itself as “a national network of physicians dedicated to ensuring patient access to approved therapies and appropriate clinical care,” but its list of “Associate Members and Financial Supporters” contains thirty drug companies, including each of the Manufacturing Defendants named in this lawsuit. In addition, the APA’s board members include doctors who have received hundreds of thousands of dollars in payments from drug companies. As discussed above, the APA has been a vocal critic of policies restricting the flow of opioids and has supported efforts to curtail the DEA’s ability to stop suspicious orders of prescription drugs.

177. The “white paper” issued by the APA in 2013 also echoed a favorite narrative of the Manufacturing Defendants, the supposed distinction between “legitimate patients” on the one hand and “addicts” on the other, asserting that one “unintended consequence” of regulating pain medication would be that “[p]atients with legitimate medical needs feel stigmatized, treated like addicts.”⁷⁵

178. Another group utilized by the Manufacturing Defendants to encourage opioid prescribing practices, a University of Wisconsin-based organization known as the **Pain & Policy**

⁷⁴ *Clinical Guideline for the Use of Chronic Opioid Therapy in Chronic Noncancer Pain*, Am. Pain Soc’y, <http://americanpainsociety.org/uploads/education/guidelines/chronic-opioid-therapy-cnccp.pdf> (last visited Aug. 13, 2018).

⁷⁵ *Prescription Pain Medication: Preserving Patient Access While Curbing Abuse*, *supra* note 42.

1 **Studies Group**, received \$2.5 million from pharmaceutical companies to promote opioid use and
 2 discourage the passing of regulations against opioid use in medical practice. The Pain & Policy
 3 Studies Group wields considerable influence over the nation's medical schools as well as within
 4 the medical field in general.⁷⁶ Purdue was the largest contributor to the Pain & Policy Studies
 5 Group, paying approximately \$1.6 million between 1999 and 2010.⁷⁷

6 179. The **Federation of State Medical Boards** (FSMB) of the United States is a
 7 national non-profit organization that represents the seventy-state medical and osteopathic boards
 8 of the United States and its territories and co-sponsors the United States Medical Licensing
 9 Examination. Beginning in 1997, FSMB developed model policy guidelines around the treatment
 10 of pain, including opioid use. The original initiative was funded by the Robert Wood Johnson
 11 Foundation, but subsequently AAPM, APS, the University of Wisconsin Pain & Policy Studies
 12 Group, and the American Society of Law, Medicine, & Ethics all made financial contributions to
 13 the project.

14 180. FSMB's 2004 *Model Policy* encourages state medical boards "to evaluate their
 15 state pain policies, rules, and regulations to identify *any regulatory restrictions or barriers that*
 16 *may impede the effective use of opioids* to relieve pain."⁷⁸ (Emphasis added).

17 181. One of the most significant barriers to convincing doctors that opioids were safe
 18 to prescribe to their patients for long-term treatment of chronic pain was the fact that many of
 19 those patients would, in fact, become addicted to opioids. If patients began showing up at their
 20 doctors' offices with obvious signs of addiction, the doctors would, of course, become concerned
 21 and likely stop prescribing opioids. And, doctors might stop believing the Manufacturing
 22 Defendants' claims that addiction risk was low.

23 ⁷⁶ *The Role of Pharmaceutical Companies in the Opioid Epidemic*, Addictions.com,
 24 <https://www.addictions.com/opiate/the-role-of-pharmaceutical-companies-in-the-opioid-epidemic/> (last visited
 Aug. 13, 2018).

25 ⁷⁷ John Fauber, *UW group ends drug firm funds*, Journal Sentinel (Apr. 20, 2011),
<http://archive.jsonline.com/watchdog/watchdogreports/120331689.html>.

26 ⁷⁸ *Model Policy for the Use of Controlled Substances for the Treatment of Pain*, Fed'n of St. Med. Boards of the
 U.S., Inc. (May 2004), <http://www.painpolicy.wisc.edu/sites/www.painpolicy.wisc.edu/files/model04.pdf>.

182. To overcome this hurdle, the Manufacturing Defendants promoted a concept called “pseudoaddiction.” These Defendants told doctors that when their patients appeared to be addicted to opioids—for example, asking for more and higher doses of opioids, increasing doses themselves, or claiming to have lost prescriptions in order to get more opioids—this was not actual addiction. Rather, the Manufacturing Defendants told doctors what appeared to be classic signs of addiction were actually just signs of undertreated pain. The solution to this “pseudoaddiction”: more opioids. Instead of warning doctors of the risk of addiction and helping patients to wean themselves off of powerful opioids and deal with their actual addiction, the Manufacturing Defendants pushed even more dangerous drugs onto patients.

183. The FSMB’s *Model Policy* gave a scientific veneer to this fictional and overstated concept. The policy defines “pseudoaddiction” as “[t]he iatrogenic syndrome resulting from the misinterpretation of relief seeking behaviors as though they are drug-seeking behaviors that are commonly seen with addiction” and states that these behaviors “resolve upon institution of effective analgesic therapy.”⁷⁹

184. In May 2012, Senate Finance Committee Chairman Max Baucus and senior Committee member Chuck Grassley initiated an investigation into the connections of the Manufacturing Defendants with medical groups and physicians who have advocated increased opioid use.⁸⁰ In addition to Purdue, Endo, and Janssen, the senators sent letters to APF, APS, AAPM, FSMB, the University of Wisconsin Pain & Policy Studies Group, the Joint Commission on Accreditation of Healthcare Organization, and the Center for Practical Bioethics, requesting from each “a detailed account of all payments/transfers received from corporations and any

⁷⁹ *Id.*

⁸⁰ Baucus, Grassley Seek Answers about Opioid Manufacturers’ Ties to Medical Groups, U.S. Senate Comm. on Fin. (May 8, 2012), <https://www.finance.senate.gov/chairmans-news/baucus-grassley-seek-answers-about-opioid-manufacturers-ties-to-medical-groups>.

related corporate entities and individuals that develop, manufacture, produce, market, or promote the use of opioid-based drugs from 1997 to the present.”⁸¹

185. On the same day as the senators’ investigation began, APF announced that it would “cease to exist, effective immediately.”⁸²

3. “It was pseudoscience”: the Manufacturing Defendants paid prominent physicians to promote their products.

186. The Manufacturing Defendants retained highly credentialed medical professionals to promote the purported benefits and minimal risks of opioids. Known as “Key Opinion Leaders” or “KOLs,” these medical professionals were often integrally involved with the front groups described above. The Manufacturing Defendants paid these KOLs substantial amounts to present at Continuing Medical Education (“CME”) seminars and conferences, and to serve on their advisory boards and on the boards of the various front groups.

187. The Manufacturing Defendants also identified doctors to serve as speakers or attend all-expense-paid trips to programs with speakers.⁸³ The Manufacturing Defendants used these trips and programs—many of them lavish affairs—to incentivize the use of opioids while downplaying their risks, bombarding doctors with messages about the safety and efficacy of opioids for treating long-term pain. Although often couched in scientific certainty, the Manufacturing Defendants’ messages were false and misleading, and helped to ensure that millions of Americans would be exposed to the profound risks of these drugs.

188. It is well documented that this type of pharmaceutical company symposium influences physicians’ prescribing, even though physicians who attend such symposia believe

⁸¹ Letter from U.S. Senate Comm. on Fin. to Am. Pain Found. (May 8, 2012), <https://www.finance.senate.gov/imo/media/doc/05092012%20Baucus%20Grassley%20Opioid%20Investigation%20Letter%20to%20American%20Pain%20Foundation2.pdf>.

⁸² Charles Ornstein and Tracy Weber, *American Pain Foundation Shuts Down as Senators Launch Investigation of Prescription Narcotics*, ProPublica (May 8, 2012, 8:57pm), <https://www.propublica.org/article/senate-panel-investigates-drug-company-ties-to-pain-groups>.

⁸³ Van Zee, *The Promotion and Marketing of OxyContin*, *supra* note 65.

that such enticements do not alter their prescribing patterns.⁸⁴ For example, doctors who were invited to these all-expenses-paid weekends in resort locations like Boca Raton, Florida, and Scottsdale, Arizona, wrote twice as many prescriptions as those who did not attend.⁸⁵

189. The KOLs gave the impression they were independent sources of unbiased information, while touting the benefits of opioids through their presentations, articles, and books. KOLs also served on committees and helped develop guidelines such as the 2009 Guidelines described above that strongly encouraged the use of opioids to treat chronic pain.

190. One of the most prominent KOLs for the Manufacturing Defendants' opioids was Dr. Russell Portenoy. A respected leader in the field of pain treatment, Dr. Portenoy was highly influential. Dr. Andrew Kolodny, cofounder of Physicians for Responsible Opioid Prescribing, described him "lecturing around the country as a religious-like figure. The megaphone for Portenoy is Purdue, which flies in people to resorts to hear him speak. It was a compelling message: 'Docs have been letting patients suffer; nobody really gets addicted; it's been studied.'"⁸⁶

191. As one organizer of CME seminars, who worked with Portenoy and Purdue, pointed out, "had Portenoy not had Purdue's money behind him, he would have published some papers, made some speeches, and his influence would have been minor. With Purdue's millions behind him, his message, which dovetailed with their marketing plans, was hugely magnified."⁸⁷

192. In recent years, some of the Manufacturing Defendants' KOLs have conceded that many of their past claims in support of opioid use lacked evidence or support in the scientific literature.⁸⁸ Dr. Portenoy himself specifically admitted that he overstated the drugs' benefits and

⁸⁴ *Id.*

⁸⁵ Harriet Ryan, Lisa Girion and Scott Glover, *OxyContin goes global* — "We're only just getting started", Los Angeles Times (Dec. 18, 2016), <http://www.latimes.com/projects/la-me-oxycontin-part3/>.

⁸⁶ Quinones, *supra* note 43, at 314.

⁸⁷ *Id.* at 136.

⁸⁸ See, e.g., John Fauber, *Painkiller boom fueled by networking*, Journal Sentinel (Feb. 18, 2012), <http://archive.jsonline.com/watchdog/watchdogreports/painkiller-boom-fueled-by-networking-dp3p2rn-139609053.html/> (finding that a key Endo KOL acknowledged that opioid marketing went too far).

1 glossed over their risks, and that he “gave innumerable lectures in the late 1980s and ‘90s about
 2 addiction that weren’t true.”⁸⁹ He mused, “Did I teach about pain management, specifically about
 3 opioid therapy, in a way that reflects misinformation? Well, against the standards of 2012, I
 4 guess I did . . . We didn’t know then what we know now.”⁹⁰

5 193. Dr. Portenoy did not need “the standards of 2012” to discern evidence-based
 6 science from baseless claims, however. When interviewed by journalist Barry Meier for his 2003
 7 book, *Pain Killer*, Dr. Portenoy was more direct: “It was pseudoscience. I guess I’m going to
 8 have always to live with that one.”⁹¹

9 194. Dr. Portenoy was perhaps the most prominent KOL for prescription opioids, but
 10 he was far from the only one. In fact, Dr. Portenoy and a doctor named Perry Fine co-wrote A
 11 *Clinical Guide to Opioid Analgesia*, which contained statements that conflict with the CDC’s
 12 2016 *Guideline for Prescribing Opioids for Chronic Pain*, such as the following examples
 13 regarding respiratory depression and addiction:

14 At clinically appropriate doses, . . . respiratory rate typically does not decline.
 15 Tolerance to the respiratory effects usually develops quickly, and doses can be
 steadily increased without risk.

16 Overall, the literature provides evidence that the outcomes of drug abuse and
 17 addiction are rare among patients who receive opioids for a short period (ie, for
 18 acute pain) and among those with no history of abuse who receive long-term
 therapy for medical indications.⁹²

19 195. Dr. Fine is a Professor of Anesthesiology at the University of Utah School of
 20 Medicine’s Pain Research Center. He has served on Purdue’s advisory board, provided medical
 21 legal consulting for Janssen, and participated in CME activities for Endo, along with serving in
 22 these capacities for several other drug companies. He co-chaired the APS-AAPM Opioid

23
 24 ⁸⁹ Thomas Catan and Evan Perez, *A Pain-Drug Champion Has Second Thoughts*, Wall Street Journal (Dec. 17,
 2012, 11:36am), <https://www.wsj.com/articles/SB10001424127887324478304578173342657044604>.

25 ⁹⁰ *Id.*

⁹¹ Meier, *supra* note 15, at 277.

26 ⁹² Perry G. Fine, MD and Russell K. Portenoy, MD, *A Clinical Guide to Opioid Analgesia* 20 and 34, McGraw-Hill
 Companies (2004), <http://www.thblack.com/links/RSD/OpioidHandbook.pdf>.

Guideline Panel, served as treasurer of the AAPM from 2007 to 2010 and as president of that group from 2011 to 2013, and was also on the board of directors of APF.⁹³

196. In 2011, he and Dr. Scott Fishman, discussed below, published a letter in *JAMA* called “Reducing Opioid Abuse and Diversion,” which emphasized the importance of maintaining patient access to opioids.⁹⁴ The editors of *JAMA* found that both doctors had provided incomplete financial disclosures and made them submit corrections listing all of their ties to the prescription painkiller industry.⁹⁵

197. Dr. Fine also failed to provide full disclosures as required by his employer, the University of Utah. For example, Dr. Fine told the university that he had received under \$5,000 in 2010 from Johnson & Johnson for providing “educational” services, but Johnson & Johnson’s website states that the company paid him \$32,017 for consulting, promotional talks, meals and travel that year.⁹⁶

198. In 2012, along with other KOLs, Dr. Fine was investigated for his ties to drug companies as part of the Senate investigation of front groups described above. When Marianne Skolek, a reporter for the online news outlet Salem-News.com and a critic of opioid overuse, wrote an article about him and another KOL being investigated, Dr. Fine fired back, sending a letter to her editor accusing her of poor journalism and saying that she had lost whatever credibility she may have had. He criticized her for linking him to Purdue, writing, “I have never had anything to do with Oxycontin development, sales, marketing or promotion; I have never

⁹³ Scott M. Fishman, MD, *Incomplete Financial Disclosures in a Letter on Reducing Opioid Abuse and Diversion*, 306 (13) JAMA 1445 (Sept. 20, 2011), <https://jamanetwork.com/journals/jama/article-abstract/1104464?redirect=true>.

⁹⁴ Perry G. Fine, MD and Scott M. Fishman, MD, *Reducing Opioid Abuse and Diversion*, 306 (4) JAMA 381 (July 27, 2011), <https://jamanetwork.com/journals/jama/article-abstract/1104144?redirect=true>.

⁹⁵ *Incomplete Financial Disclosures in: Reducing Opioid Abuse and Diversion*, 306 (13) JAMA 1446 (Oct. 5, 2011), <https://jamanetwork.com/journals/jama/fullarticle/1104453>.

⁹⁶ Weber and Ornstein, *Two Leaders in Pain Treatment*, *supra* note 73.

1 been a Purdue Pharma speaker”—neglecting to mention, of course, that he served on Purdue’s
2 advisory board, as the *JAMA* editors had previously forced him to disclose.⁹⁷

3 199. Another Utah physician, Dr. Lynn Webster, was the director of Lifetree Clinical
4 Research & Pain Clinic in Salt Lake City from 1990 to 2010, and in 2013 was the president of
5 AAPM (one of the front groups discussed above). Dr. Webster developed a five-question survey
6 he called the Opioid Risk Tool, which he asserted would “predict accurately which individuals
7 may develop aberrant behaviors when prescribed opioids for chronic pain.”⁹⁸ He published
8 books titled *The Painful Truth: What Chronic Pain Is Really Like and Why It Matters to Each of*
9 *Us* and *Avoiding Opioid Abuse While Managing Pain*.

10 200. Dr. Webster and the Lifetree Clinic were investigated by the DEA for
11 overprescribing opioids after twenty patients died from overdoses. In keeping with the opioid
12 industry’s promotional messages, Dr. Webster apparently believed the solution to patients’
13 tolerance or addictive behaviors was more opioids: he prescribed staggering quantities of pills.
14 Tina Webb, a Lifetree patient who overdosed in 2007, was taking as many as thirty-two pain
15 pills a day in the year before she died, all while under doctor supervision.⁹⁹ Carol Ann Bosley,
16 who sought treatment for pain at Lifetree after a serious car accident and multiple spine
17 surgeries, quickly became addicted to opioids and was prescribed increasing quantities of pills; at
18 the time of her death, she was on seven different medications totaling approximately 600 pills a
19 month.¹⁰⁰ Another woman, who sought treatment from Lifetree for chronic low back pain and
20
21

22 ⁹⁷ Marianne Skolek, *Doctor Under Senate Investigation Lashes Out at Journalist*, Salem News (Aug. 12, 2012,
23 8:45pm), <http://www.salem-news.com/articles/august122012/perry-fine-folo-ms.php>.

24 ⁹⁸ Lynn Webster and RM Webster, *Predicting aberrant behaviors in opioid-treated patients: preliminary validation*
25 *of the Opioid Risk Tool* 6 (6) Pain Med. 432 (Nov.-Dec. 2005), <https://www.ncbi.nlm.nih.gov/pubmed/16336480>.

26 ⁹⁹ Jesse Hyde and Daphne Chen, *The untold story of how Utah doctors and Big Pharma helped drive the national*
opioid epidemic, Deseret News (Oct. 26, 2017, 12:01am), <https://www.deseretnews.com/article/900002328/the-untold-story-of-how-utah-doctors-and-big-pharma-helped-drive-the-national-opioid-epidemic.html>.

¹⁰⁰ Stephanie Smith, *Prominent pain doctor investigated by DEA after patient deaths*, CNN (Dec. 20, 2013,
7:06am), <http://www.cnn.com/2013/12/20/health/pain-pillar/index.html>.

1 headaches, died at age forty-two after Lifetree clinicians increased her prescriptions to fourteen
2 different drugs, including multiple opioids, for a total of 1,158 pills a month.¹⁰¹

3 201. By these numbers, Lifetree resembles the pill mills and “bad actors” that the
4 Manufacturing Defendants blame for opioid overuse. But Dr. Webster was an integral part of
5 Defendants’ marketing campaigns, a respected pain specialist who authored numerous CMEs
6 sponsored by Endo and Purdue. And the Manufacturing Defendants promoted his Opioid Risk
7 Tool and similar screening questionnaires as measures that allow powerful opioids to be
8 prescribed for chronic pain.

9 202. Even in the face of patients’ deaths, Dr. Webster continues to promote a pro-
10 opioid agenda, even asserting that alternatives to opioids are risky because “[i]t’s not hard to
11 overdose on NSAIDs or acetaminophen.”¹⁰² He argued on his website in 2015 that DEA
12 restrictions on the accessibility of hydrocodone harm patients, and in 2017 tweeted in response to
13 CVS Caremark’s announcement that it will limit opioid prescriptions that “CVS Caremark’s new
14 opioid policy is wrong, and it won’t stop illegal drugs.”¹⁰³

15 203. Another prominent KOL is Dr. Scott M. Fishman, the Chief of the Department of
16 Pain Medicine at University of California, Davis. He has served as president of APF and AAPM,
17 and as a consultant and a speaker for Purdue, in addition to providing the company grant and
18 research support. He also has had financial relationships with Endo and Janssen. He wrote a
19 book for the FSMB called *Responsible Opioid Use: A Physician’s Guide*, which was distributed
20 to over 165,000 physicians in the U.S.

21 204. Dr. Fishman and Dr. Fine, along with Dr. Seddon Savage, published an editorial
22 in the Seattle Times in 2010, arguing that Washington legislation proposed to combat

23 ¹⁰¹ *Id.*

24 ¹⁰² APF releases opioid medication safety module, Drug Topics (May 10, 2011),
25 [http://drugtopics.modernmedicine.com/drug-topics/news/modernmedicine/modern-medicine-news/apf-releases-
opioid-medication-safety-module](http://drugtopics.modernmedicine.com/drug-topics/news/modernmedicine/modern-medicine-news/apf-releases-opioid-medication-safety-module).

26 ¹⁰³ Lynn Webster, MD (@LynnRWebsterMD), Twitter (Dec. 7, 2017, 5:45pm),
<https://twitter.com/LynnRWebsterMD/status/938887130545360898>.

1 prescription opioid abuse would harm patients, in particular by requiring chronic pain patients to
 2 consult with a pain specialist before receiving a prescription for a moderate to high dose of an
 3 opioid.¹⁰⁴

4 205. These KOLs and others—respected specialists in pain medicine—proved to be
 5 highly effective spokespeople for the Manufacturing Defendants.

6 **4. The Manufacturing Defendants used “unbranded” advertising as a platform**
 7 **for their misrepresentations about opioids.**

8 206. The Manufacturing Defendants also aggressively promoted opioids through
 9 “unbranded advertising” to generally tout the benefits of opioids without specifically naming a
 10 particular brand-name opioid drug. Instead, unbranded advertising is usually framed as “disease
 11 awareness”—encouraging consumers to “talk to your doctor” about a certain health condition
 12 without promoting a specific product. A trick often used by pharmaceutical companies,
 13 unbranded advertising gives the pharmaceutical companies considerable leeway to make
 14 sweeping claims about health conditions or classes of drugs. In contrast, a “branded”
 15 advertisement that identifies a specific medication and its indication (i.e., the condition which the
 16 drug is approved to treat) must also include possible side effects and contraindications—what the
 17 FDA Guidance on pharmaceutical advertising refers to as “fair balance.” Branded advertising is
 18 also subject to FDA review for consistency with the drug’s FDA-approved label.

19 207. Unbranded advertising allows pharmaceutical manufacturers to sidestep those
 20 requirements; “fair balance” and consistency with a drug’s label are not required.

21 208. By engaging in unbranded advertising, the Manufacturing Defendants were and
 22 are able to avoid FDA review and issue general statements to the public including that opioids
 23 improve function, that addiction usually does not occur, and that withdrawal can easily be
 24

25 ¹⁰⁴ Perry G. Fine, Scott M. Fishman, and Seddon R. Savage, *Bill to combat prescription abuse really will harm*
 26 *patients in pain*, Seattle Times (Mar. 16, 2010, 4:39pm),
http://old.seattletimes.com/html/opinion/2011361572_guest17fine.html.

1 managed. The Manufacturing Defendants' unbranded advertisements either did not disclose the
2 risks of addiction, abuse, misuse, and overdose, or affirmatively denied or minimized those risks.

3 209. Through the various marketing channels described above—all of which the
4 Manufacturing Defendants controlled, funded, and facilitated, and for which they are legally
5 responsible—these Defendants made false or misleading statements about opioids despite the
6 lack of scientific evidence to support their claims, while omitting the true risk of addiction and
7 death.

8 **D. Specific Misrepresentations Made by the Manufacturing Defendants.**

9 210. All the Manufacturing Defendants have made and/or continue to make false or
10 misleading claims in the following areas: (1) the low risk of addiction to opioids, (2) opioids'
11 efficacy for chronic pain and ability to improve patients' quality of life with long-term use, (3)
12 the lack of risk associated with higher dosages of opioids, (4) the need to prescribe more opioids
13 to treat withdrawal symptoms, and (5) that risk-mitigation strategies and abuse-deterrent
14 technologies allow doctors to safely prescribe opioids for chronic use. These illustrative but non-
15 exhaustive categories of the Manufacturing Defendants' misrepresentations about opioids are
16 described in detail below.

17 **1. The Manufacturing Defendants falsely claimed that the risk of opioid abuse**
18 **and addiction was low.**

19 211. Collectively, the Manufacturing Defendants have made a series of false and
20 misleading statements about the low risk of addiction to opioids over the past twenty years. The
21 Manufacturing Defendants have also failed to take sufficient remedial measures to correct their
22 false and misleading statements.

23 212. The Manufacturing Defendants knew that many physicians were hesitant to
24 prescribe opioids other than for acute or cancer-related pain because of concerns about addiction.
25 Because of this general perception, sales messaging about the low risk of addiction was a
26 fundamental prerequisite misrepresentation.

213. Purdue launched OxyContin in 1996 with the statement that OxyContin's patented continuous-release mechanism "is believed to reduce the abuse liability." This statement, which appeared in OxyContin's label and which sales representatives were taught to repeat verbatim, was unsupported by any studies, and was patently false. The continuous-release mechanism was simple to override, and the drug correspondingly easy to abuse. This fact was known, or should have been known, to Purdue prior to its launch of OxyContin, because people had been circumventing the same continuous-release mechanism for years with MS Contin, which in fact commanded a high street price because of the dose of pure narcotic it delivered. In addition, with respect to OxyContin, Purdue researchers notified company executives, including Raymond and Richard Sackler, by email that patients in their clinical trials were abusing the drug despite the timed-release mechanism.¹⁰⁵

214. In 2007, as noted above, Purdue pleaded guilty to misbranding a drug, a felony under the Food, Drug, and Cosmetic Act. 21 U.S.C. § 331(a)(2). As part of its guilty plea, Purdue agreed that certain Purdue supervisors and employees had, "with the intent to defraud or mislead, marketed and promoted OxyContin as less addictive, less subject to abuse and diversion, and less likely to cause tolerance and withdrawal than other pain medications" in the following ways:

Trained PURDUE sales representatives and told some health care providers that it was more difficult to extract the oxycodone from an OxyContin tablet for the purpose of intravenous abuse, although PURDUE's own study showed that a drug abuser could extract approximately 68% of the oxycodone from a single 10mg OxyContin tablet by crushing the tablet, stirring it in water, and drawing the solution through cotton into a syringe;

Told PURDUE sales representatives they could tell health care providers that OxyContin potentially creates less chance for addiction than immediate-release opioids;

¹⁰⁵ WBUR On Point interview, *supra* note 21.

1 Sponsored training that taught PURDUE sales supervisors that OxyContin had
 2 fewer “peak and trough” blood level effects than immediate-release opioids
 resulting in less euphoria and less potential for abuse than short-acting opioids;

3 Told certain health care providers that patients could stop therapy abruptly without
 4 experiencing withdrawal symptoms and that patients who took OxyContin would
 not develop tolerance to the drug; and

5 Told certain health care providers that OxyContin did not cause a “buzz” or
 6 euphoria, caused less euphoria, had less addiction potential, had less abuse
 7 potential, was less likely to be diverted than immediate-release opioids, and could
 be used to “weed out” addicts and drug seekers.¹⁰⁶

8 215. All of these statements were false and misleading. But Purdue had not stopped
 9 there. Purdue—and later the other Defendants—manipulated scientific research and utilized
 10 respected physicians as paid spokespeople to convey its misrepresentations about low addiction
 11 risk in much more subtle and pervasive ways, so that the idea that opioids used for chronic pain
 12 posed a low addiction risk became so widely accepted in the medical community that Defendants
 13 were able to continue selling prescription opioids for chronic pain—even after Purdue’s criminal
 14 prosecution.

15 216. When it launched OxyContin, Purdue knew it would need data to overcome
 16 decades of wariness regarding opioid use. It needed some sort of research to back up its
 17 messaging. But Purdue had not conducted any studies about abuse potential or addiction risk as
 18 part of its application for FDA approval for OxyContin. Purdue (and, later, the other Defendants)
 19 found this “research” in the form of a one-paragraph letter to the editor published in the *New*
 20 *England Journal of Medicine* (NEJM) in 1980.

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 26 ¹⁰⁶ *United States v. Purdue Frederick Co.*, *supra* note 25; *see also*, Plea Agreement, *United States v. Purdue Frederick Co.*, No. 1:07-cr-00029 (W.D. Va. May 10, 2007).

217. This letter, by Dr. Hershel Jick and Jane Porter, declared the incidence of addiction “rare” for patients treated with opioids.¹⁰⁷ They had analyzed a database of hospitalized patients who were given opioids in a controlled setting to ease suffering from acute pain. These patients were not given long-term opioid prescriptions or provided opioids to administer to themselves at home, nor was it known how frequently or infrequently and in what doses the patients were given their narcotics. Rather, it appears the patients were treated with opioids for short periods of time under in-hospital doctor supervision.

**ADDICTION RARE IN PATIENTS TREATED
WITH NARCOTICS**

To the Editor: Recently, we examined our current files to determine the incidence of narcotic addiction in 39,946 hospitalized medical patients¹ who were monitored consecutively. Although there were 11,882 patients who received at least one narcotic preparation, there were only four cases of reasonably well documented addiction in patients who had no history of addiction. The addiction was considered major in only one instance. The drugs implicated were meperidine in two patients,² Percodan in one, and hydromorphone in one. We conclude that despite widespread use of narcotic drugs in hospitals, the development of addiction is rare in medical patients with no history of addiction.

JANE PORTER
HERSHEL JICK, M.D.
Boston Collaborative Drug
Surveillance Program

Waltham, MA 02154 Boston University Medical Center

1. Jick H, Miettinen OS, Shapiro S, Lewis GP, Siskind Y, Slone D. Comprehensive drug surveillance. JAMA. 1970; 213:1455-60.
2. Miller RR, Jick H. Clinical effects of meperidine in hospitalized medical patients. J Clin Pharmacol. 1978; 18:180-8.

218. As Dr. Jick explained to a journalist years later, he submitted the statistics to NEJM as a letter because the data were not robust enough to be published as a study, and that one could not conclude anything about long-term use of opioids from his figures.¹⁰⁸ Dr. Jick also recalled that no one from drug companies or patient advocacy groups contacted him for more information about the data.¹⁰⁹

¹⁰⁷ Jane Porter and Herschel Jick, MD, *Addiction Rare in Patients Treated with Narcotics*, 302(2) N Engl J Med. 123 (Jan. 10, 1980), <http://www.nejm.org/doi/pdf/10.1056/NEJM198001103020221>.

¹⁰⁸ Meier, *supra* note 15, at 174.

¹⁰⁹ *Id.*

219. Nonetheless, the Manufacturing Defendants regularly invoked this letter as proof of the low addiction risk in connection with taking opioids despite its obvious shortcomings. These Defendants' egregious misrepresentations based on this letter included claims that *less than one percent* of opioid users become addicted.

220. The limited facts of the study did not deter the Manufacturing Defendants from using it as definitive proof of opioids' safety. The enormous impact of the Manufacturing Defendants' misleading amplification of this letter was well documented in another letter published in NEJM on June 1, 2017, describing the way the one-paragraph 1980 letter had been irresponsibly cited and in some cases "grossly misrepresented." In particular, the authors of this letter explained:

[W]e found that a five-sentence letter published in the *Journal* in 1980 was heavily and uncritically cited as evidence that addiction was rare with long-term opioid therapy. We believe that this citation pattern contributed to the North American opioid crisis by helping to shape a narrative that allayed prescribers' concerns about the risk of addiction associated with long-term opioid therapy . . .¹¹⁰

221. Unfortunately, by the time of this analysis and the CDC's findings in 2016, the damage had already been done. "It's difficult to overstate the role of this letter," said Dr. David Juurlink of the University of Toronto, who led the analysis. "It was the key bit of literature that helped the opiate manufacturers convince front-line doctors that addiction is not a concern."¹¹¹

¹¹⁰ Pamela T.M. Leung, B.Sc. Pharm., Erin M. Macdonald, M.Sc., Matthew B. Stanbrook, M.D., Ph.D., Irfan Al Dhalla, M.D., David N. Juurlink, M.D., Ph.D., *A 1980 Letter on the Risk of Opioid Addiction*, 376 N Engl J Med 2194-95 (June 1, 2017), <http://www.nejm.org/doi/full/10.1056/NEJMc1700150#t=article>.

¹¹¹ *Painful words: How a 1980 letter fueled the opioid epidemic*, STAT News (May 31, 2017), <https://www.statnews.com/2017/05/31/opioid-epidemic-nejm-letter/>.

222. The Manufacturing Defendants successfully manipulated the 1980 Porter and Jick letter as the “evidence” supporting their fundamental misrepresentation that the risk of opioid addiction was low when opioids were prescribed to treat pain. For example, in its 1996 press release announcing the release of OxyContin, Purdue advertised that the “fear of addiction is exaggerated” and quoted the chairman of the American Pain Society Quality of Care Committee, who claimed that “there is very little risk of addiction from the proper uses of these [opioid] drugs for pain relief.”¹¹²

PR Newswire

May 31, 1996, Friday - 15:47 Eastern Time

NEW HOPE FOR MILLIONS OF AMERICANS SUFFERING FROM PERSISTENT

The fear of addiction is exaggerated.

One cause of patient resistance to appropriate pain treatment – the fear of addiction – is largely unfounded. According to Dr. Max, “Experts agree that most pain caused by surgery or cancer can be relieved, primarily by carefully adjusting the dose of opioid (narcotic) pain reliever to each patient's need, and that there is very little risk of addiction from the proper uses of these drugs for pain relief.”

Paul D. Goldenheim, M.D., Vice President of **Purdue Pharma** L.P. in Norwalk, Connecticut, agrees with this assessment. “Proper use of medication is an essential weapon in the battle against persistent pain. But too often fear, misinformation and poor communication stand in the way of their legitimate use.”

223. Dr. Portenoy, the Purdue KOL mentioned previously, also stated in a promotional video from the 1990s that “the likelihood that the treatment of pain using an opioid drug which is prescribed by a doctor will lead to addiction is extremely low.”¹¹³

¹¹² Press Release, OxyContin, *New Hope for Millions of Americans Suffering from Persistent Pain: Long-Acting OxyContin Tablets Now Available to Relieve Pain* (May 31, 1996, 3:47pm), <http://documents.latimes.com/oxycontin-press-release-1996/>.

¹¹³ Catan and Perez, *supra* note 89.



224. Purdue also specifically used the Porter and Jick letter in its 1998 promotional video, “I got my life back,” in which Dr. Alan Spanos says, “In fact, the rate of addiction amongst pain patients who are treated by doctors is *much less than 1%*.”¹¹⁴



225. The Porter and Jick letter was also used on Purdue’s “Partners Against Pain” website, which was available in the early 2000s, where Purdue claimed that the addiction risk with OxyContin was very low.¹¹⁵

¹¹⁴ Our Amazing World, *Purdue Pharma OxyContin Commercial*, <https://www.youtube.com/watch?v=Er78Dj5hyeI> (last visited Aug. 13, 2018) (emphasis added).

¹¹⁵ Van Zee, *The Promotion and Marketing of OxyContin*, *supra* note 65.

226. The Porter and Jick letter was used frequently in literature given to prescribing physicians and to patients who were prescribed OxyContin.¹¹⁶

227. In addition to the Porter and Jick letter, the Manufacturing Defendants exaggerated the significance of a study published in 1986 regarding cancer patients treated with opioids. Conducted by Dr. Portenoy and another pain specialist, Dr. Kathleen Foley, the study involved only 38 patients, who were treated for non-malignant cancer pain with low doses of opioids (the majority were given less than 20 MME/day, the equivalent of only 13 mg of oxycodone).¹¹⁷ Of these thirty-eight patients, only two developed problems with opioid abuse, and Dr. Portenoy and Dr. Foley concluded that “opioid maintenance therapy can be a safe, salutary and more humane alternative to the options of surgery or no treatment in those patients with intractable non-malignant pain and no history of drug abuse.”¹¹⁸ Notwithstanding the small sample size, low doses of opioids involved, and the fact that all the patients were cancer patients, the Manufacturing Defendants used this study as “evidence” that high doses of opioids were safe for the treatment of chronic non-cancer pain.

228. The Manufacturing Defendants’ repeated misrepresentations about the low risk of opioid addiction were so effective that this concept became part of the conventional wisdom. Dr. Nathaniel Katz, a pain specialist, recalls learning in medical school that previous fears about addiction were misguided, and that doctors should feel free to allow their patients the pain relief that opioids can provide. He did not question this until one of his patients died from an overdose. Then, he searched the medical literature for evidence of the safety and efficacy of opioid treatment for chronic pain. “There’s not a shred of research on the issue. All these so-called

¹¹⁶ Art Van Zee, M.D., *The OxyContin Abuse Problem: Spotlight on Purdue Pharma’s Marketing* (Aug. 22, 2001), <https://web.archive.org/web/20170212210143/https://www.fda.gov/ohrms/dockets/dockets/01n0256/c000297-A.pdf>.

¹¹⁷ Russell K. Portenoy and Kathleen M. Foley, *Chronic Use of Opioid Analgesics in Non-Malignant Pain: Report of 38 Cases*, 25 *Pain* 171-86 (1986), <https://www.ncbi.nlm.nih.gov/pubmed/2873550>.

¹¹⁸ *Id.*

1 experts in pain are dedicated and have been training me that opioids aren't as addictive as we
2 thought. But what is that based on? It was based on nothing."¹¹⁹

3 229. At a hearing before the House of Representatives' Subcommittee on Oversight
4 and Investigations of the Committee on Energy and Commerce in August 2001, Purdue
5 continued to emphasize "legitimate" treatment, dismissing cases of overdose and death as
6 something that would not befall "legitimate" patients: "Virtually all of these reports involve
7 people who are abusing the medication, not patients with legitimate medical needs under the
8 treatment of a healthcare professional."¹²⁰

9 230. Purdue spun this baseless "legitimate use" distinction out even further in a patient
10 brochure about OxyContin, called "A Guide to Your New Pain Medicine and How to Become a
11 Partner Against Pain." In response to the question, "Aren't opioid pain medications like
12 OxyContin Tablets 'addicting'? Even my family is concerned about this," Purdue claimed that
13 there was no need to worry about addiction if taking opioids for legitimate, "medical" purposes:

14 Drug addiction means using a drug to get "high" rather than to relieve pain. You
15 are taking opioid pain medication for medical purposes. The medical purposes are
16 clear and the effects are beneficial, not harmful.

17 231. Similarly, Dr. David Haddox, Senior Medical Director for Purdue, cavalierly
18 stated, "[w]hen this medicine is used appropriately to treat pain under a doctor's care, it is not
19 only effective, it is safe."¹²¹ He went so far as to compare OxyContin to celery, because even
20 celery would be harmful if injected: "If I gave you a stalk of celery and you ate that, it would be
21 healthy for you. But if you put it in a blender and tried to shoot it into your veins, it would not be
22 good."¹²²

23 ¹¹⁹ Quinones, *supra* note 43, at 188-89.

24 ¹²⁰ *Oxycontin: Its Use and Abuse: Hearing Before the H. Subcomm. on Oversight and Investigations of the Comm.*
25 *on Energy and Commerce*, 107th Cong. 1 (Aug. 28, 2001) (statement of Michael Friedman, Executive Vice
26 President, Chief Operating Officer, Purdue Pharma, L.P.), <https://www.gpo.gov/fdsys/pkg/CHRG-107hhrg75754/html/CHRG-107hhrg75754.htm>.

¹²¹ Roger Alford, *Deadly OxyContin abuse expected to spread in the U.S.*, Charleston Gazette, Feb. 9, 2001.

¹²² *Id.*

232. Purdue sales representatives also repeated these misstatements regarding the low risk for addiction to doctors across the country.¹²³ Its sales representatives targeted primary care physicians in particular, downplaying the risk of addiction and, as one doctor observed, “promot[ing] among primary care physicians a more liberal use of opioids.”¹²⁴

233. Purdue sales representatives were instructed to “distinguish between iatrogenic addiction (<1% of patients) and substance abusers/diversion (about 10% of the population abuse something: weed; cocaine; heroin; alcohol; valium; etc.).”¹²⁵

234. Purdue also marketed OxyContin for a wide variety of conditions and to doctors who were not adequately trained in pain management.¹²⁶

235. As of 2003, Purdue’s Patient Information guide for OxyContin contained the following language regarding addiction:

Concerns about abuse, addiction, and diversion should not prevent the proper management of pain. The development of addiction to opioid analgesics in properly managed patients with pain has been reported to be rare. However, data are not available to establish the true incidence of addiction in chronic pain patients.

236. Although Purdue has acknowledged it has made some misrepresentations about the safety of its opioids,¹²⁷ it has done nothing to address the ongoing harms of their misrepresentations; in fact, it continues to make those misrepresentations today.

237. Defendant Endo also made dubious claims about the low risk of addiction. For instance, it sponsored a website, PainKnowledge.com, on which in 2009 it claimed that “[p]eople

¹²³ Barry Meier, *In Guilty Plea, OxyContin Maker to Pay \$600 Million*, New York Times (May 10, 2007), <http://www.nytimes.com/2007/05/10/business/11drug-web.html>.

¹²⁴ Van Zee, *The Promotion and Marketing of OxyContin*, *supra* note 65.

¹²⁵ Meier, *supra* note 15, at 269.

¹²⁶ *OxyContin Abuse and Diversion and Efforts to Address the Problem*, *supra* note 30.

¹²⁷ Following the conviction in 2007 of three of its executives for misbranding OxyContin, Purdue released a statement in which they acknowledged their false statements. “Nearly six years and longer ago, some employees made, or told other employees to make, certain statements about OxyContin to some health care professionals that were inconsistent with the F.D.A.-approved prescribing information for OxyContin and the express warnings it contained about risks associated with the medicine. The statements also violated written company policies requiring adherence to the prescribing information.”

1 who take opioids as prescribed usually do not become addicted.”¹²⁸ The website has since been
2 taken down.

3 238. In another website, PainAction.com—which is still currently available today—
4 Endo also claimed that “most chronic pain patients do not become addicted to the opioid
5 medications that are prescribed for them.”¹²⁹

6 239. In a pamphlet titled “Understanding Your Pain: Taking Oral Opioid Analgesics,”
7 Endo assured patients that addiction is something that happens to people who take opioids for
8 reasons other than pain relief, “such as unbearable emotional problems”¹³⁰:

9
10 Some questions you may have are:

11 *Is it wrong to take opioids for pain?*

- 12 ♦ No. Pain relief is an important medical
13 reason to take opioids as prescribed
14 by your doctor. Addicts take opioids
15 for other reasons, such as unbearable
16 emotional problems. Taking opioids as
17 prescribed for pain relief is not addiction.

18 *How can I be sure I’m not addicted?*

- 19 ♦ Addiction to an opioid would mean that
20 your pain has gone away but you still
21 take the medicine regularly when you
22 don’t need it for pain, maybe just to
23 escape from your problems.
- 24 ♦ Ask yourself: Would I want to take this
25 medicine if my pain went away? If you
26 answer no, you are taking opioids for
the right reasons—to relieve your pain
and improve your function. You are not
addicted.

128 German Lopez, *The growing number of lawsuits against opioid companies, explained*, Vox (Feb. 27, 2018, 2:25pm), <https://www.vox.com/policy-and-politics/2017/6/7/15724054/opioid-companies-epidemic-lawsuits>.

129 *Opioid medication and addiction*, Pain Action (Aug. 17, 2017), <https://www.painaction.com/opioid-medication-addiction/>.

130 *Understanding Your Pain: Taking Oral Opioid Analgesics*, Endo Pharms. (2004), http://www.thblack.com/links/RSD/Understand_Pain_Opioid_Analgesics.pdf.

240. In addition, Endo made statements in pamphlets and publications that most health care providers who treat people with pain agree that most people do not develop an addiction problem. These statements also appeared on websites sponsored by Endo, such as Opana.com.

241. In its currently active website, PrescribeResponsibly.com, Defendant Janssen states that concerns about opioid addiction are “overestimated” and that “true addiction occurs only in a small percentage of patients.”¹³¹

Use of Opioid Analgesics in Pain Management



Other Opioid Analgesic Concerns

Aside from medical issues related to opioid analgesics, there are nonmedical issues that may have an impact on prescribing patterns and patient use of these drugs. Practitioners are often concerned about prescribing opioid analgesics due to potential legal issues and questions of addiction.^{15,16} By the same token, patients report similar concerns about developing an addiction to opioid analgesics.¹⁷ While these concerns are not without some merit, it would appear that they are often overestimated. According to clinical opinion polls, true addiction occurs only in a small percentage of patients with chronic pain who receive chronic opioid analgesic therapy.¹⁸



¹³¹ Keith Candiotti, M.D., *Use of Opioid Analgesics in Pain Management*, Prescribe Responsibly, <http://www.prescriberesponsibly.com/articles/opioid-pain-management> (last modified July 2, 2015).

242. Similarly, in a 2009 patient education video titled “Finding Relief: Pain Management for Older Adults,” Janssen sponsored a video by the American Academy of Pain Medicine that indicated that opioids are rarely addictive. The video has since been taken down.¹³²

243. Janssen also approved and distributed a patient education guide in 2009 that attempted to counter the “myth” that opioids are addictive, claiming that “[m]any studies show that opioids are rarely addictive when used properly for the management of chronic pain.”¹³³

244. In addition, all the Manufacturing Defendants used third parties and front groups to further their false and misleading statements about the safety of opioids.

245. For example, in testimony for the Hearing to Examine the Effects of the Painkiller OxyContin, Focusing on Risks and Benefits, in front of the Senate Health, Education, Labor and Pensions Committee in February 2002, Dr. John D. Giglio, Executive Director of the APF, the organization which, as described above, received the majority of its funding from opioid manufacturers, including Purdue, stated that “opioids are safe and effective, and only in rare cases lead to addiction.”¹³⁴ Along with Dr. Giglio’s testimony, the APF submitted a short background sheet on “the scope of the undertreatment of pain in the U.S.,” which asserted that “opioids are often the best” treatment for pain that hasn’t responded to other techniques, but that patients and many doctors “lack even basic knowledge about these options and fear that powerful pain drugs will [c]ause addiction.” According to the APF, “most studies show that less than 1% of patients become addicted, which is medically different from becoming physically dependent.”¹³⁵

¹³² Molly Huff, *Finding Relief: Pain Management for Older Adults*, Ctrs. for Pain Mgmt. (Mar. 9, 2011), <http://www.managepaintoday.com/news/-Finding-Relief-Pain-Management-for-Older-Adults>.

¹³³ Lopez, *supra* note 128.

¹³⁴ *Oxycontin: Balancing Risks and Benefits: Hearing of the S. Comm. on Health, Education, Labor and Pensions*, 107th Cong. 2 (Feb. 12, 2002) (testimony of John D. Giglio, M.A., J.D., Executive Director, American Pain Foundation), <https://www.help.senate.gov/imo/media/doc/Giglio.pdf>.

¹³⁵ *Id.*

246. The APF further backed up Purdue in an amicus curiae brief filed in an Ohio appeals court in December 2002, in which it claimed that “medical leaders have come to understand that the small risk of abuse does not justify the withholding of these highly effective analgesics from chronic pain patients.”¹³⁶

247. In a 2007 publication titled “Treatment Options: A Guide for People Living with Pain,” APF downplayed the risk of addiction and argued that concern about this risk should not prevent people from taking opioids: “Restricting access to the most effective medications for treating pain is not the solution to drug abuse or addiction.”¹³⁷ APF also tried to normalize the dangers of opioids by listing opioids as one of several “[c]ommon drugs that can cause physical dependence,” including steroids, certain heart medications, and caffeine.¹³⁸

248. The Manufacturing Defendants’ repeated statements about the low risk of addiction when taking opioids as prescribed for chronic pain were blatantly false and were made with reckless disregard for the potential consequences.

2. The Manufacturing Defendants falsely claimed that opioids were proven effective for chronic pain and would improve quality of life.

249. Not only did the Manufacturing Defendants falsely claim that the risk of addiction to prescription opioids was low, these Defendants represented that there was a significant upside to long-term opioid use, including that opioids could restore function and improve quality of life.¹³⁹

¹³⁶ Brief Amici Curiae of American Pain Foundation, National Foundation for the Treatment of Pain, and The Ohio Pain Initiative, in Support of Defendants/Appellants, *Howland v. Purdue Pharma, L.P.*, Appeal No. CA 2002 09 0220 (Butler Co., Ohio 12th Court of Appeals, Dec. 23, 2002), <https://ia801005.us.archive.org/23/items/279014-howland-apf-amicus/279014-howland-apf-amicus.pdf>.

¹³⁷ *Treatment Options: A Guide for People Living with Pain*, Am. Pain Found., <https://assets.documentcloud.org/documents/277605/apf-treatmentoptions.pdf> (last visited Aug. 13, 2018).

¹³⁸ *Id.*

¹³⁹ This case *does not* request or require the Court to specifically adjudicate whether opioids are appropriate for the treatment of chronic, non-cancer pain—though the scientific evidence strongly suggests they are not.

250. Such claims were viewed as a critical part of the Manufacturing Defendants' marketing strategies. For example, an internal Purdue report from 2001 noted the lack of data supporting improvement in quality of life with OxyContin treatment:

Janssen has been stressing decreased side effects, especially constipation, as well as patient quality of life, as supported by patient rating compared to sustained release morphine . . . We do not have such data to support OxyContin promotion. . . In addition, Janssen has been using the "life uninterrupted" message in promotion of Duragesic for non-cancer pain, stressing that Duragesic "helps patients think less about their pain." This is a competitive advantage based on our inability to make any quality of life claims.¹⁴⁰

251. Despite the lack of data supporting improvement in quality of life, Purdue ran a full-page ad for OxyContin in the Journal of the American Medical Association in 2002, proclaiming, "There Can Be Life With Relief," and showing a man happily fly-fishing alongside his grandson.¹⁴¹ This ad earned a warning letter from the FDA, which admonished, "It is particularly disturbing that your November ad would tout 'Life With Relief' yet fail to warn that patients can die from taking OxyContin."¹⁴²

252. Purdue also consistently tried to steer any concern away from addiction and focus on its false claims that opioids were effective and safe for treating chronic pain. At a hearing before the House of Representatives' Subcommittee on Oversight and Investigations of the Committee on Energy and Commerce in August 2001, Michael Friedman, Executive Vice President and Chief Operating Officer of Purdue, testified that "even the most vocal critics of opioid therapy concede the value of OxyContin in the legitimate treatment of pain," and that "OxyContin has proven itself an effective weapon in the fight against pain, returning many patients to their families, to their work, and to their ability to enjoy life."¹⁴³

¹⁴⁰ Meier, *supra* note 15, at 281.

¹⁴¹ *Id.* at 280.

¹⁴² Chris Adams, *FDA Orders Purdue Pharma To Pull Its OxyContin Ads*, Wall Street Journal (Jan. 23, 2003, 12:01am), <https://www.wsj.com/articles/SB1043259665976915824>.

¹⁴³ *Oxycontin: Its Use and Abuse*, *supra* note 120.

1 253. Purdue sponsored the development and distribution of an APF guide in 2011
 2 which claimed that “multiple clinical studies have shown that opioids are effective in improving
 3 daily function, psychological health, and health-related quality of life for chronic pain patients.”
 4 This guide is still available today.

5 254. Purdue also ran a series of advertisements of OxyContin in 2012 in medical
 6 journals titled “Pain vignettes,” which were styled as case studies of patients with persistent pain
 7 conditions and for whom OxyContin was recommended to improve their function.

8 255. Purdue and Endo also sponsored and distributed a book in 2007 to promote the
 9 claim that pain relief from opioids, by itself, improved patients’ function. The book remains for
 10 sale online today.

11 256. Endo’s advertisements for Opana ER claimed that use of the drug for chronic pain
 12 allowed patients to perform demanding tasks like construction and portrayed Opana ER users as
 13 healthy and unimpaired.

14 257. Endo’s National Initiative on Pain Control (NIPC) website also claimed in 2009
 15 that with opioids, “your level of function should improve; you may find you are now able to
 16 participate in activities of daily living, such as work and hobbies, that you were not able to enjoy
 17 when your pain was worse.”

18 258. Endo further sponsored a series of CME programs through NIPC which claimed
 19 that chronic opioid therapy has been “shown to reduce pain and depressive symptoms and
 20 cognitive functioning.”

21 259. Through PainKnowledge.org, Endo also supported and sponsored guidelines that
 22 stated, among other things, that “Opioid Medications are a powerful and often highly effective
 23 tool in treating pain,” and that “they can help restore comfort, function, and quality of life.”¹⁴⁴
 24

25 ¹⁴⁴*Informed Consent for Using Opioids to Treat Pain*, Painknowledge.org (2007),
 26 [https://www.mainequalitycounts.org/image_upload/Opioid%20Informed%20Consent%20Formatted_1_23_2008.p
 df.](https://www.mainequalitycounts.org/image_upload/Opioid%20Informed%20Consent%20Formatted_1_23_2008.pdf)

1 260. In addition, Janssen sponsored and edited patient guides which stated that
2 “opioids may make it easier for people to live normally.” The guides listed expected functional
3 improvements from opioid use, including sleeping through the night, and returning to work,
4 recreation, sex, walking, and climbing stairs.

5 261. Janssen also sponsored, funded, and edited a website which featured an interview
6 edited by Janssen that described how opioids allowed a patient to “continue to function.” This
7 video is still available today.

8 262. Furthermore, sales representatives for the Manufacturing Defendants
9 communicated and continue to communicate the message that opioids will improve patients’
10 function, without appropriate disclaimers.

11 263. The Manufacturing Defendants’ statements regarding opioids’ ability to improve
12 function and quality of life are false and misleading. As the CDC’s *Guideline for Prescribing*
13 *Opioids for Chronic Pain* (the “2016 CDC Guideline” or “Guideline”)¹⁴⁵ confirms, not a single
14 study supports these claims.

15 264. In fact, to date, there have been no long-term studies that demonstrate that opioids
16 are effective for treating long-term or chronic pain. Instead, reliable sources of information,
17 including from the CDC in 2016, indicate that there is “[n]o evidence” to show “a long-term
18 benefit of opioids in pain and function versus no opioids for chronic pain.”¹⁴⁶ By contrast,
19 significant research has demonstrated the colossal dangers of opioids. The CDC, for example,
20 concluded that “[e]xtensive evidence shows the possible harms of opioids (including opioid use
21 disorder, overdose, and motor vehicle injury)” and that “[o]pioid pain medication use presents
22 serious risks, including overdose and opioid use disorder.”¹⁴⁷

23
24
25 ¹⁴⁵ 2016 CDC Guideline, *supra* note 31.

26 ¹⁴⁶ *Id.*

¹⁴⁷ *Id.*

1 **3. The Manufacturing Defendants falsely claimed doctors and patients could**
 2 **increase opioid usage indefinitely without added risk.**

3 265. The Manufacturing Defendants also made false and misleading statements
 4 claiming that there is no dosage ceiling for opioid treatment. These misrepresentations were
 5 integral to the Manufacturing Defendants' promotion of prescription opioids for two reasons.
 6 First, the idea that there was no upward limit was necessary for the overarching deception that
 7 opioids are appropriate treatment for chronic pain. As discussed above, people develop a
 8 tolerance to opioids' analgesic effects, so that achieving long-term pain relief requires constantly
 9 increasing the dose. Second, the dosing misrepresentation was necessary for the claim that
 10 OxyContin and competitor drugs allowed 12-hour dosing.

11 266. Twelve-hour dosing is a significant marketing advantage for any medication,
 12 because patient compliance is improved when a medication only needs to be taken twice a day.
 13 For prescription painkillers, the 12-hour dosing is even more significant because shorter-acting
 14 painkillers did not allow patients to get a full night's sleep before the medication wore off. A
 15 Purdue memo to the OxyContin launch team stated that "OxyContin's positioning statement is
 16 'all of the analgesic efficacy of immediate-release oxycodone, with convenient q12h dosing,'" and further that "[t]he convenience of q12h dosing was emphasized as the most important
 17 benefit."¹⁴⁸

18
 19 267. Purdue executives therefore maintained the messaging of 12-hour dosing even
 20 when many reports surfaced that OxyContin did not last 12 hours. Instead of acknowledging a
 21 need for more frequent dosing, Purdue instructed its representatives to push higher-strength pills.

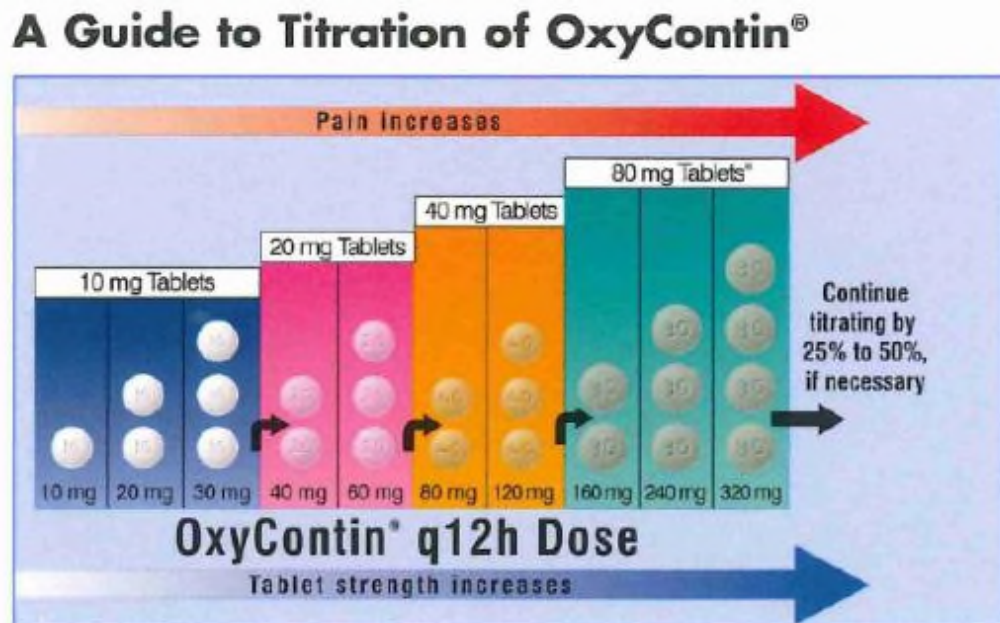
22 268. For example, in a 1996 sales strategy memo from a Purdue regional manager, the
 23 manager emphasized that representatives should "convinc[e] the physician that there is no need"
 24 for prescribing OxyContin in shorter intervals than the recommended 12-hour interval, and
 25 instead the solution is prescribing higher doses. The manager directed representatives to discuss

26 ¹⁴⁸ *OxyContin launch*, Los Angeles Times (May 5, 2016), <http://documents.latimes.com/oxycontin-launch-1995/>.

with physicians that there is “no[] upward limit” for dosing and ask “if there are any reservations in using a dose of 240mg-320mg of OxyContin.”¹⁴⁹

269. As doctors began prescribing OxyContin at shorter intervals in the late 1990s, Purdue directed its sales representatives to “refocus” physicians on 12-hour dosing. One sales manager instructed her team that anything shorter “needs to be nipped in the bud. NOW!!”¹⁵⁰

270. These misrepresentations were incredibly dangerous. As noted above, opioid dosages at or above 50 MME/day double the risk of overdose compared to 20 MME/day, and 50 MME is equal to just 33 mg of oxycodone. Notwithstanding the risks, Purdue’s 2003 Conversion Guide for OxyContin contained the following diagram for increasing dosage up to 320 mg:



271. In a 2004 response letter to the FDA, Purdue tried to address concerns that patients who took OxyContin more frequently than 12 hours would be at greater risk of side effects or adverse reactions. Purdue contended that the peak plasma concentrations of oxycodone would not increase with more frequent dosing, and therefore no adjustments to the package

¹⁴⁹ Sales manager on 12-hour dosing, Los Angeles Times (May 5, 2016), <http://documents.latimes.com/sales-manager-on12-hour-dosing-1996/>.

¹⁵⁰ Harriet Ryan, Lisa Girion, and Scott Glover, ‘You Want a Description of Hell?’ OxyContin’s 12-Hour Problem (May 5, 2016), <http://www.latimes.com/projects/oxycontin-part1/>.

1 labeling or 12-hour dosing regimen were needed.¹⁵¹ But these claims were false, and Purdue's
 2 suggestion that there was no upper limit or risk associated with increased dosage was incredibly
 3 misleading.

4 272. Suggesting that it recognized the danger of its misrepresentations of no dose
 5 ceiling, Purdue discontinued the OxyContin 160 mg tablet in 2007 and stated that this step was
 6 taken "to reduce the risk of overdose accompanying the abuse of this dosage strength."¹⁵²

7 273. But still Purdue and the other Manufacturing Defendants worked hard to protect
 8 their story. In March 2007, Dr. Gary Franklin, Medical Director for the Washington State
 9 Department of Labor & Industries, published the *Interagency Guideline on Opioid Dosing for*
 10 *Chronic Non-Cancer Pain*. Developed in collaboration with providers in Washington State who
 11 had extensive experience in the evaluation and treatment of patients with chronic pain, the
 12 guideline recommended a maximum daily dose of opioids to protect patients.

13 274. In response, Purdue sent correspondence to Dr. Franklin specifically indicating,
 14 among other things, that "limiting access to opioids for persons with chronic pain is not the
 15 answer" and that the "safety and efficacy of OxyContin doses greater than 40 mg every 12 hours
 16 in patients with chronic nonmalignant pain" was well established. Purdue even went so far as to
 17 represent to Dr. Franklin that even if opioid treatment produces significant adverse effects in a
 18 patient, "this does not preclude a trial of another opioid."

19 275. In 2010, Purdue published a Risk Evaluation and Mitigation Strategy ("REMS")
 20 for OxyContin, but even the REMS does not address concerns with increasing dosage, and
 21 instead advises prescribers that "dose adjustments may be made every 1-2 days"; "it is most
 22 appropriate to increase the q12h dose"; the "total daily dose can usually be increased by 25% to
 23

24 ¹⁵¹ *Purdue Response to FDA, 2004*, Los Angeles Times (May 5, 2016), [http://documents.latimes.com/purdue-](http://documents.latimes.com/purdue-response-fda-2004/)
 25 [response-fda-2004/](http://documents.latimes.com/purdue-response-fda-2004/).

26 ¹⁵² *OxyContin Tablets Risk Management Program*, Purdue Pharma L.P.,
[https://web.archive.org/web/20170215064438/https://www.fda.gov/ohrms/dockets/DOCKETS/07p0232/07p-0232-](https://web.archive.org/web/20170215064438/https://www.fda.gov/ohrms/dockets/DOCKETS/07p0232/07p-0232-cp00001-03-Exhibit-02-Part-1-vol1.pdf)
[cp00001-03-Exhibit-02-Part-1-vol1.pdf](https://web.archive.org/web/20170215064438/https://www.fda.gov/ohrms/dockets/DOCKETS/07p0232/07p-0232-cp00001-03-Exhibit-02-Part-1-vol1.pdf) (revised May 18, 2007).

50%”; and if “significant adverse reactions occur, treat them aggressively until they are under control, then resume upward titration.”¹⁵³

276. In 2012, APF claimed on its website that there was no “ceiling dose” for opioids for chronic pain.¹⁵⁴ APF also made this claim in a guide sponsored by Purdue, which is still available online.

277. Accordingly, Purdue continued to represent both publicly and privately that increased opioid usage was safe and did not present additional risk at higher doses.

278. Janssen also made the same misrepresentations regarding the disadvantages of dosage limits for other pain medicines in a 2009 patient education guide, while failing to address the risks of dosage increases with opioids.

279. Endo, on a website it sponsors, PainKnowledge.com, also made the claim in 2009 that opioid dosages could be increased indefinitely.

280. In the “Understanding Your Pain” pamphlet discussed above, Endo assures opioid users that concern about developing tolerance to the drugs’ pain-relieving effect is “not a problem,” and that “[t]he dose can be increased” and “[y]ou won’t ‘run out’ of pain relief.”¹⁵⁵

¹⁵³ *OxyContin Risk Evaluation and Mitigation Strategy*, Purdue Pharma L.P., <https://web.archive.org/web/20170215190303/https://www.fda.gov/downloads/Drugs/DrugSafety/PostmarketDrugSafetyInformationforPatientsandProviders/UCM220990.pdf> (last modified Nov. 2010).

¹⁵⁴ Noah Nesin, M.D., FAAFP, *Responsible Opioid Prescribing*, PCHC https://www.mainequalitycounts.org/image_upload/Keynote-%20Managing%20Chronic%20Pain%20and%20Opioids_Nesin.pdf (last visited Aug. 13, 2018).

¹⁵⁵ *Understanding Your Pain: Taking Oral Opioid Analgesics*, *supra* note 130.

ENDO
PHARMACEUTICALS
Toll Free: 800-452-3636
Web: www.endo.com

Understanding Your Pain

Taking Oral Opioid Analgesics

This brochure was developed by Margo McCallery, RN, MS, FAAN, and Chris Parsons, RN, MS, FAAN authors of *Pain: Clinical Manual* (2nd ed. Mosby: 1999), edited by Russell K. Portney, MD.

© 2004 Pain Management 1 NUP135

(Have you felt over the last week?)

- Addiction to an opioid would mean that your pain has gone away but you still take the medicine regularly when you don't need it for pain, maybe just to escape from your problems.
- Ask yourself: Would I want to take the medicine if my pain went away? If you answer no, you are taking opioids for the right reasons—to relieve your pain and improve your function. You are not addicted.

IF I TAKE THIS OPIOID NOW, WILL IT WORK LATER WHEN I REALLY NEED IT?

Some patients with chronic pain worry about this, but it is not a problem.

- The dose can be increased or other medicines can be added.
- The opioid "breaks" if you need it.

WHAT CAN I DO ABOUT SIDE EFFECTS?

Talk to your doctor, nurse, or pharmacist about the side effects of opioids. If they occur, remember that most opioid side effects can be treated or prevented.

Constipation

- Constipation from opioids is very common, but it can be prevented. If it does occur, it can be treated.
- Prevention is the best approach. If you take opioids daily, you need to eat more fiber and drink more liquids than you usually do. Many people also need to take a laxative. The most common type is a combination of stool softener and mild stimulant laxative. These can be purchased without a prescription include Peri-Colace® capsules or syrup and Senokot® tablets. Ask your pharmacist about less expensive generic forms.

Nausea or vomiting (stomach upset)

- This does not always occur, but if it does, it can be treated. Ask your doctor, nurse, or pharmacist for medicine to relieve this. After a few days, the nausea usually stops.
- Try eating small and breathing slowly through your mouth.
- Nausea medicines that you can buy without a prescription include Dramamine® tablets and Enamox® oral solution.
- If your pain is under good control, you may be able to reduce the nausea by taking a lower dose of opioid.

Drowsiness (sleepiness)

- Some degree of sleepiness would be normal when you start taking an opioid, but after a few days the drowsiness usually goes away.

281. Dosage limits with respect to opioids are particularly important not only because of the risk of addiction but also because of the potentially fatal side effect of respiratory depression. Endo's "Understanding Your Pain" pamphlet minimized this serious side effect, calling it "slowed breathing," declaring that it is "very rare" when opioids are used "appropriately," and never stating that it could be fatal:

"Slowed breathing"

- ◆ The medical term for "slowed breathing" is "respiratory depression."
- ◆ This is very rare when oral opioids are used appropriately for pain relief.
- ◆ If you become so sleepy that you cannot make yourself stay awake, you may be in danger of slowed breathing. Stop taking your opioid and call your doctor immediately.

1 **4. The Manufacturing Defendants falsely instructed doctors and patients that**
 2 **more opioids were the solution when patients presented symptoms of**
 3 **addiction.**

4 282. Not only did the Manufacturing Defendants hide the serious risks of addiction
 5 associated with opioids, they actively worked to prevent doctors from taking steps to prevent or
 6 address opioid addiction in their patients.

7 283. One way that the Manufacturing Defendants worked to obstruct appropriate
 8 responses to opioid addiction was to push a concept called “pseudoaddiction.” Dr. David
 9 Haddox—who later became a Senior Medical Director for Purdue—published a study in 1989
 10 coining the term, which he characterized as “the iatrogenic syndrome of abnormal behavior
 11 developing as a direct consequence of inadequate pain management.”¹⁵⁶ (“Iatrogenic” describes a
 12 condition induced by medical treatment.) In other words, he claimed that people on prescription
 13 opioids who exhibited classic signs of addiction—“abnormal behavior”—were not addicted, but
 14 rather simply suffering from under-treatment of their pain. His solution for pseudoaddiction?
 15 More opioids.

16 284. Although this concept was formed based on a single case study, it proved to be a
 17 favorite trope in the Manufacturing Defendants’ marketing schemes. For example, using this
 18 study, Purdue informed doctors and patients that signs of addiction are actually the signs of
 19 under-treated pain which should be treated with even more opioids. Purdue reassured doctors and
 20 patients, telling them that “chronic pain has been historically undertreated.”¹⁵⁷

21 285. The Manufacturing Defendants continued to spread the concept of
 22 pseudoaddiction through the APF, which even went so far as to compare opioid addicts to coffee
 23 drinkers. In a 2002 court filing, APF wrote that “[m]any pain patients (like daily coffee drinkers)
 24 claim they are ‘addicted’ when they experience withdrawal symptoms associated with physical

25 ¹⁵⁶ David E. Weissman and J. David Haddox, *Opioid pseudoaddiction--an iatrogenic syndrome*, 36(3) Pain 363-66
 26 (Mar. 1989), <https://www.ncbi.nlm.nih.gov/pubmed/2710565>.

¹⁵⁷ *Oxycontin: Its Use and Abuse*, *supra* note 120.

dependence as they decrease their dose. But unlike actual addicts, such individuals, if they resume their opioid use, will only take enough medication to alleviate their pain . . .”¹⁵⁸

286. In a 2007 publication titled “Treatment Options: A Guide for People Living with Pain,” the APF claimed: “*Physical dependence is normal*; any patient who is taking an opioid on a regular basis for a few days should be assumed to be physically dependent. This does **NOT** mean you are addicted.”¹⁵⁹ In this same publication, the APF asserted that “people who are not substance abusers” may also engage in “unacceptable” behaviors such as “increasing the dose without permission or obtaining the opioid from multiple sources,” but that such behaviors do not indicate addiction and instead reflect a “desire to obtain pain relief.”¹⁶⁰

287. Purdue published a REMS for OxyContin in 2010, and in the associated Healthcare Provider Training Guide stated that “[b]ehaviors that suggest drug abuse exist on a continuum, and pain-relief seeking behavior can be mistaken for drug-seeking behavior.”¹⁶¹

288. Purdue worked, and continues to work, to create confusion about what addiction is. For example, Purdue continues to emphasize that abuse and addiction are separate and distinct from physical dependence. Regardless of whether these statements may be technically correct, they continue to add ambiguity over the risks and benefits of opioids.

289. Endo sponsored an NIPC CME program in 2009 which promoted the concept of pseudoaddiction by teaching that a patient’s aberrant behavior was the result of untreated pain. Endo substantially controlled NIPC by funding its projects, developing content, and reviewing NIPC materials.

¹⁵⁸ APF Brief Amici Curiae, *supra* note 136, at 10-11.

¹⁵⁹ *Treatment Options: A Guide for People Living with Pain*, *supra* note 137.

¹⁶⁰ *Id.*

¹⁶¹ *OxyContin Risk Evaluation and Mitigation Strategy*, *supra* note 153.

1 290. A 2001 paper which was authored by a doctor affiliated with Janssen stated that
 2 “[m]any patients presenting to a doctor’s office asking for pain medications are accused of drug
 3 seeking. In reality, most of these patients may be undertreated for their pain syndrome.”¹⁶²

4 291. In 2009, on a website it sponsored, Janssen stated that pseudoaddiction is different
 5 from true addiction “because such behaviors can be resolved with effective pain
 6 management.”¹⁶³

7 292. Indeed, on its currently active website PrescribeResponsibly.com, Janssen defines
 8 pseudoaddiction as “a syndrome that causes patients to seek additional medications due to
 9 inadequate pharmacotherapy being prescribed. Typically, when the pain is treated appropriately,
 10 the inappropriate behavior ceases.”¹⁶⁴

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22 ¹⁶² Howard A. Heit, MD, FACP, FASAM, *The truth about pain management: the difference between a pain patient
 and an addicted patient*, 5 *European Journal of Pain* 27-29 (2001),
 23 <http://www.med.uottawa.ca/courses/totalpain/pdf/doc-34.pdf>.

24 ¹⁶³ Chris Morran, *Ohio: Makers Of OxyContin, Percocet & Other Opioids Helped Fuel Drug Epidemic By
 Misleading Doctors, Patients*, *Consumerist* (May 31, 2017, 2:05pm), [https://consumerist.com/2017/05/31/ohio-
 makers-of-oxycontin-percocet-other-opioids-helped-fuel-drug-epidemic-by-misleading-doctors-patients/](https://consumerist.com/2017/05/31/ohio-makers-of-oxycontin-percocet-other-opioids-helped-fuel-drug-epidemic-by-misleading-doctors-patients/).

25 ¹⁶⁴ Howard A. Heit, MD, FACP, FASAM and Douglas L. Gourlay, MD, MSc, FRCPC, FASAM, *What a Prescriber
 Should Know Before Writing the First Prescription, Prescribe Responsibly*,
 26 <http://www.prescriberesponsibly.com/articles/before-prescribing-opioids#pseudoaddiction> (last modified July 2,
 2015).

What a Prescriber Should Know Before Writing the First Prescription



TABLE 1: Definitions

8. **Pseudoaddiction** is a syndrome that causes patients to seek additional medications due to inadequate pharmacotherapy being prescribed. Typically when the pain is treated appropriately, the inappropriate behavior ceases.²⁵



293. As set forth in more detail below, these statements were false and misleading as evidenced by, *inter alia*, the findings made by the CDC in 2016. Indeed, there is simply no evidence that pseudoaddiction is a real phenomenon. As research compiled by the CDC and others makes clear, pseudoaddiction is pseudoscience—nothing more than a concept Defendants seized upon to help sell more of their actually addicting drugs.

5. **The Manufacturing Defendants falsely claimed that risk-mitigation strategies, including tapering and abuse-deterrent technologies, made it safe to prescribe opioids for chronic use.**

294. Even when the Manufacturing Defendants acknowledge that opioids pose some risk of addiction, they dismiss these concerns by claiming that addiction can be easily avoided and addressed through simple steps. In order to make prescribers feel more comfortable about

1 starting patients on opioids, the Manufacturing Defendants falsely communicated to doctors that
 2 certain screening tools would allow them to reliably identify patients at higher risk of addiction
 3 and safely prescribe opioids, and that tapering the dose would be sufficient to manage cessation
 4 of opioid treatment. Both assertions are false.

5 295. For instance, as noted above, Purdue published a REMS for OxyContin in 2010,
 6 in which it described certain steps that needed to be followed for safe opioid use. Purdue stressed
 7 that all patients should be screened for their risk of abuse or addiction, and that such screening
 8 could curb the incidence of addiction.¹⁶⁵

9 296. The APF also proclaimed in a 2007 booklet, sponsored in part by Purdue, that
 10 “[p]eople with the disease of addiction may abuse their medications, engaging in unacceptable
 11 behaviors like increasing the dose without permission or obtaining the opioid from multiple
 12 sources, among other things. Opioids get into the hands of drug dealers and persons with an
 13 addictive disease as a result of pharmacy theft, forged prescriptions, Internet sales, and even
 14 from other people with pain. It is a problem in our society that needs to be addressed through
 15 many different approaches.”¹⁶⁶

16 297. On its current website for OxyContin,¹⁶⁷ Purdue acknowledges that certain
 17 patients have higher risk of opioid addiction based on history of substance abuse or mental
 18 illness—a statement which, even if accurate, obscures the significant risk of addiction for all
 19 patients, including those without such a history, and comports with statements it has recently
 20 made that it is “bad apple” patients, and not the opioids, that are arguably the source of the
 21 opioid crisis:
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 24

25 ¹⁶⁵ *Oxycontin Risk Evaluation and Mitigation Strategy*, *supra* note 153.

26 ¹⁶⁶ *Treatment Options: A Guide for People Living with Pain*, *supra* note 137.

¹⁶⁷ OxyContin, <https://www.oxycontin.com/index.html> (last visited Aug. 13, 2018).

Assess each patient's risk for opioid addiction, abuse, or misuse prior to prescribing OxyContin, and monitor all patients receiving OxyContin for the development of these behaviors and conditions. Risks are increased in patients with a personal or family history of substance abuse (including drug or alcohol abuse or addiction) or mental illness (e.g., major depression). The potential for these risks should not, however, prevent the proper management of pain in any given patient. Patients at increased risk may be prescribed opioids such as OxyContin, but use in such patients necessitates intensive counseling about the risks and proper use of OxyContin along with intensive monitoring for signs of addiction, abuse, and misuse.

298. Additionally, on its current website, Purdue refers to publicly available tools that can assist with prescribing compliance, such as patient-prescriber agreements and risk assessments.¹⁶⁸

299. Purdue continues to downplay the severity of addiction and withdrawal and claims that dependence can easily be overcome by strategies such as adhering to a tapering schedule to successfully stop opioid treatment. On the current website for OxyContin, it instructs that “[w]hen discontinuing OxyContin, gradually taper the dosage. Do not abruptly discontinue OxyContin.”¹⁶⁹ And on the current OxyContin Medication Guide, Purdue also states that one should “taper the dosage gradually.”¹⁷⁰ As a general matter, tapering is a sensible strategy for cessation of treatment with a variety of medications, such as steroids or antidepressants. But the suggestion that tapering is sufficient in the context of chronic use of potent opioids is misleading and dangerous, and sets patients up for withdrawal and addiction.

¹⁶⁸ *ER/LA Opioid Analgesics REMS*, Purdue, <http://www.purduepharma.com/healthcare-professionals/responsible-use-of-opioids/remis/> (last visited Aug. 13, 2018).

¹⁶⁹ Oxycontin.com, *supra* note 167.

¹⁷⁰ *OxyContin Full Prescribing Information*, Purdue Pharma LP, <http://app.purduepharma.com/xmlpublishing/pi.aspx?id=o> (last visited Aug. 13, 2018).

1 300. In its “Dear Healthcare Professional” letter in 2010, Purdue instructed doctors to
2 gradually taper someone off OxyContin to prevent signs and symptoms of withdrawal in patients
3 who were physically dependent.¹⁷¹ Nowhere does Purdue warn doctors or patients that tapering
4 may be inadequate to safely end opioid treatment and avoid addiction.

5 301. Other Manufacturing Defendants make similar claims. For instance, Endo
6 suggests that risk-mitigation strategies enable the safe prescription of opioids. In its currently
7 active website, Opana.com, Endo states that assessment tools should be used to assess addiction
8 risk, but that “[t]he potential for these risks should not, however, prevent proper management of
9 pain in any given patient.”¹⁷²

10 302. On the same website, Endo makes similar statements about tapering, stating
11 “[w]hen discontinuing OPANA ER, gradually taper the dosage.”¹⁷³

12 303. Janssen also states on its currently active website, PrescribeResponsibly.com, that
13 the risk of opioid addiction “can usually be managed” through tools such as “opioid agreements”
14 between patients and doctors.¹⁷⁴

15 304. Each Manufacturing Defendant’s statements about tapering misleadingly implied
16 that gradual tapering would be sufficient to alleviate any risk of withdrawal or addiction while
17 taking opioids.

18 305. The Manufacturing Defendants have also made and continue to make false and
19 misleading statements about the purported abuse-deterrent properties of their opioid pills to
20 suggest these reformulated pills are not susceptible to abuse. In so doing, the Manufacturing
21 Defendants have increased their profits by selling more pills for substantially higher prices.

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25 ¹⁷¹ *OxyContin Risk Evaluation and Mitigation Strategy*, *supra* note 153.

26 ¹⁷² Opana ER, Endo Pharmaceuticals, Inc., <http://www.opana.com> (last visited Aug. 13, 2018).

¹⁷³ *Id.*

¹⁷⁴ Heit & Gourlay, *supra* note 164.

1 306. For instance, since at least 2001, Purdue has contended that “abuse resistant
2 products can reduce the incidence of abuse.”¹⁷⁵ Until recently, Purdue’s website touted abuse-
3 deterrent properties by saying they “can make a difference.”¹⁷⁶

4 307. On August 17, 2015, Purdue announced the launch of a new website, “Team
5 Against Opioid Abuse,” which it said was “designed to help healthcare professionals and
6 laypeople alike learn about different abuse-deterrent technologies and how they can help in the
7 reduction of misuse and abuse of opioids.”¹⁷⁷ This website appears to no longer be active.

8 308. A 2013 study which was authored by at least two doctors who at one time
9 worked for Purdue stated that “[a]buse-deterrent formulations of opioid analgesics can reduce
10 abuse.”¹⁷⁸ In another study from 2016 with at least one Purdue doctor as an author, the authors
11 claimed that abuse decreased by as much as 99% in some situations after abuse-deterrent
12 formulations were introduced.¹⁷⁹

13 309. Interestingly, one report found that the original safety label for OxyContin, which
14 instructed patients not to crush the tablets because it would have a rapid release effect, may have
15 inadvertently given opioid users ideas for techniques to get high from these drugs.¹⁸⁰
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19 ¹⁷⁵ *Oxycontin: Its Use and Abuse*, *supra* note 120.

20 ¹⁷⁶ *Opioids with Abuse-Deterrent Properties*, Purdue, [http://www.purduepharma.com/healthcare-
21 professionals/responsible-use-of-opioids/opioids-with-abuse-deterrent-properties/](http://www.purduepharma.com/healthcare-professionals/responsible-use-of-opioids/opioids-with-abuse-deterrent-properties/) (last visited May 16, 2018); see
22 also [https://web.archive.org/web/20180302203422/http://www.purduepharma.com/healthcare-
23 professionals/responsible-use-of-opioids/opioids-with-abuse-deterrent-properties/](https://web.archive.org/web/20180302203422/http://www.purduepharma.com/healthcare-professionals/responsible-use-of-opioids/opioids-with-abuse-deterrent-properties/).

24 ¹⁷⁷ *Purdue Pharma L.P. Launches TeamAgainstOpioidAbuse.com*, Purdue (Aug. 17, 2015),
25 <http://www.purduepharma.com/news-media/2015/08/purdue-pharma-l-p-launches-teamagainstopioidabuse-com/>.

26 ¹⁷⁸ Paul M. Coplan, Hrishikesh Kale, Lauren Sandstrom, Craig Landau, and Howard D. Chilcoat, *Changes in
oxycodone and heroin exposures in the National Poison Data System after introduction of extended-release
oxycodone with abuse-deterrent characteristics*, 22 (12) *Pharmacoepidemiol Drug Saf.* 1274-82 (Sept. 30, 2013),
<https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4283730/>.

¹⁷⁹ Paul M. Coplan, Howard D. Chilcoat, Stephen Butler, Edward M. Sellers, Aditi Kadakia, Venkatesh
Harikrishnan, J. David Haddox, and Richard C. Dart, *The effect of an abuse-deterrent opioid formulation
(OxyContin) on opioid abuse-related outcomes in the postmarketing setting*, 100 *Clin. Pharmacol. Ther.* 275-86
(June 22, 2016), <http://onlinelibrary.wiley.com/doi/10.1002/cpt.390/full>.

¹⁸⁰ *OxyContin Abuse and Diversion and Efforts to Address the Problem*, *supra* note 30.

310. In 2012, Defendant Endo replaced the formula for Opana ER with a new formula with abuse-deterrent properties that it claimed would make Opana ER resistant to manipulation from users to snort or inject it. But the following year, the FDA concluded:

While there is an increased ability of the reformulated version of Opana ER to resist crushing relative to the original formulation, study data show that the reformulated version's extended-release features can be compromised when subjected to other forms of manipulation, such as cutting, grinding, or chewing, followed by swallowing.

Reformulated Opana ER can be readily prepared for injection, despite Endo's claim that these tablets have "resistance to aqueous extraction (i.e., poor syringeability)." It also appears that reformulated Opana ER can be prepared for snorting using commonly available tools and methods.

The postmarketing investigations are inconclusive, and even if one were to treat available data as a reliable indicator of abuse rates, one of these investigations also suggests the troubling possibility that a higher percentage of reformulated Opana ER abuse is via injection than was the case with the original formulation.¹⁸¹

311. Despite the FDA's determination that the evidence did not support Endo's claims of abuse-deterrence, Endo advertised its reformulated pills as "crush resistant" and directed its sales representatives to represent the same to doctors. Endo improperly marketed Opana ER as crush-resistant, when Endo's own studies showed that the pill could be crushed and ground. In 2016, Endo reached an agreement with the Attorney General of the State of New York that required Endo to discontinue making such statements.¹⁸²

312. The Manufacturing Defendants' assertions that their reformulated pills could curb abuse were false and misleading, as the CDC's 2016 Guideline, discussed below, confirm.

313. Ultimately, even if a physician prescribes opioids after screening for abuse risk, advising a patient to taper, and selecting brand-name, abuse-deterrent formulations, chronic

¹⁸¹ FDA Statement: Original Opana ER Relisting Determination, U.S. Food & Drug Admin. (May 10, 2013), <https://wayback.archive-it.org/7993/20171102214123/https://www.fda.gov/Drugs/DrugSafety/ucm351357.htm>.

¹⁸² Press Release, Attorney General Eric T. Schneiderman, A.G. Schneiderman Announces Settlement with Endo Health Solutions Inc. & Endo Pharmaceuticals Inc. Over Marketing of Prescription Opioid Drugs (Mar. 3, 2016), <https://ag.ny.gov/press-release/ag-schneiderman-announces-settlement-endo-health-solutions-inc-endo-pharmaceuticals>.

1 opioid use still comes with significant risks of addiction and abuse. The Manufacturing
 2 Defendants' statements to the contrary were designed to create a false sense of security and
 3 assure physicians that they could safely prescribe potent narcotics to their patients.

4 **E. Research by Washington State's Department of Labor and Industries Highlights the**
 5 **Falseness of the Manufacturing Defendants' Claims.**

6 314. Contrary to the Manufacturing Defendants' misrepresentations about the benefits
 7 and risks of opioids, growing evidence suggests that using opioids to treat chronic pain leads to
 8 overall negative outcomes, delaying or preventing recovery and providing little actual relief, all
 9 while presenting serious risks of overdose.

10 315. One place where this evidence surfaced is the Washington State Department of
 11 Labor and Industries ("L&I"). The Department of L&I runs the state's workers' compensation
 12 program, which covers all employees in the state, other than those who work for large companies
 13 and government entities. In 2000, L&I's new chief pharmacist, Jaymie Mai, noticed an increase
 14 in prescription of opioids for chronic pain, approximately 50 to 100 cases a month.¹⁸³ As she
 15 took a closer look at the prescription data, she discovered some of these same workers were
 16 dying from opioid overdoses. That workers suffered back pain or sprained knees on the job was
 17 nothing new, but workers dying from their pain medication was assuredly not business as usual.
 18 Mai reported what she was seeing to L&I's Medical Director, Dr. Gary Franklin.¹⁸⁴

19 316. In addition to being L&I's Medical Director, Dr. Franklin is a research professor
 20 at the University of Washington in the departments of Environmental Health, Neurology, and
 21 Health Services. Dr. Franklin and Mai undertook a thorough analysis of all recorded deaths in
 22 the state's workers' comp system. In 2005, they published their findings in the American Journal
 23 of Industrial Medicine.¹⁸⁵

24 ¹⁸³ Quinones, *supra* note 43, at 203.

25 ¹⁸⁴ *Id.*

26 ¹⁸⁵ Gary M. Franklin, M.D., MPH, Jaymie Mai, Pharm.D., Thomas Wickizer, Ph.D., Judith A. Turner, Ph.D.,
 Deborah Fulton-Kehoe, Ph.D., MPH, and Linda Grant, BSN, MBA, *Opioid dosing trends and mortality in
 Washington State Workers' Compensation, 1996-2002*, 48 Am J Ind Med 91-99 (2005).

1 317. Their research showed that the total number of opioid prescriptions paid for by
 2 the Workers' Compensation Program tripled between 1996 and 2006.¹⁸⁶ Not only did the number
 3 of prescriptions balloon, so too did the doses; from 1996 to 2002 the mean daily morphine
 4 equivalent dose ("MED") nearly doubled, and remained that way through 2006.¹⁸⁷ As injured
 5 Washington workers were given more prescriptions of higher doses of opioids, the rates of
 6 opioid overdoses among that population jumped, from zero in 1996 to more than twenty in 2005.
 7 And in 2009, over thirty people receiving opioid prescriptions through the Workers'
 8 Compensation Program died of an opioid overdose.¹⁸⁸

9 318. Armed with these alarming statistics, Dr. Franklin, in conjunction with other
 10 doctors in Washington, set out to limit the doses of opioids prescribed through the workers'
 11 compensation program. As part of that effort, in 2007 the Agency Medical Directors Group
 12 launched an Interagency Guideline on Opioid Dosing, aimed at reducing the numbers of opioid
 13 overdoses. Through this, and other related efforts, both the rates of opioid prescriptions and the
 14 sizes of doses have declined in Washington, beginning in 2009. As opioid prescriptions rates for
 15 injured workers have declined, so too has the death rate among this population.¹⁸⁹

16 319. Moreover, additional research from L&I showed that the use of opioids to treat
 17 pain after an injury actually prevents or slows a patient's recovery.

18 320. In a study of employees who had suffered a low back injury on the job, Dr.
 19 Franklin showed that if an injured worker was prescribed opioids soon after the injury, high
 20 doses of opioids, or opioids for more than a week, the employee was far more likely to
 21
 22

23 ¹⁸⁶ Gary M. Franklin, M.D., MPH, Jaymie Mai, Pharm.D., Thomas Wickizer, Ph.D., Judith Turner, Ph.D., Mark
 24 Sullivan, M.D., Ph.D., Thomas Wickizer, Ph.D., and Deborah Fulton-Kehoe, Ph.D., *Bending the Prescription*
Opioid Dosing and Mortality Curves: Impact of the Washington State Opioid Dosing Guideline, 55 Am J Ind Med
 25 325, 327 (2012).

26 ¹⁸⁷ *Id.* at 327-28.

¹⁸⁸ *Id.* at 328.

¹⁸⁹ *Id.*

1 experience negative health outcomes than the same employee who was not prescribed opioids in
2 these manners.

3 321. Specifically, the study showed that, after adjusting for the baseline covariates,
4 injured workers who received a prescription opioid for more than seven days during the first six
5 weeks after the injury were 2.2 times more likely to remain disabled a year later than workers
6 with similar injuries who received no opioids at all. Similarly, those who received two
7 prescriptions of opioids for the injury were 1.8 times more likely to remain disabled a year after
8 their injury than workers who received no opioids at all, and those receiving daily doses higher
9 than 150 MED were over twice as likely to be on disability a year later, relative to workers who
10 received no opioids.¹⁹⁰

11 322. In sum, not only do prescription opioids present significant risks of addiction and
12 overdose, but they also hinder patient recovery after an injury.

13 323. This dynamic presents problems for employers, too, who bear significant costs
14 when their employees do not recover quickly from workplace injuries. Employers are left
15 without their labor force and may be responsible for paying for the injured employee's disability
16 for long periods of time.

17 **F. The 2016 CDC Guideline and Other Recent Studies Confirm That the**
18 **Manufacturing Defendants' Statements About the Risks and Benefits of Opioids**
19 **Are Patently False.**

20 324. Contrary to the statements made by the Manufacturing Defendants in their well-
21 orchestrated campaign to tout the benefits of opioids and downplay their risks, recent studies
22 confirm the Manufacturing Defendants' statements were false and misleading.

23 325. The CDC issued its *Guideline for Prescribing Opioids for Chronic Pain* on March
24 15, 2016.¹⁹¹ The 2016 CDC Guideline, approved by the FDA, "provides recommendations for

25 ¹⁹⁰ Franklin, GM, Stover, BD, Turner, JA, Fulton-Kehoe, D, Wickizer, TM, *Early opioid prescription and*
26 *subsequent disability among workers with back injuries: the Disability Risk Identification Study Cohort*, 33 Spine
199, 201-202.

¹⁹¹ 2016 CDC Guideline, *supra* note 31.

1 primary care clinicians who are prescribing opioids for chronic pain outside of active cancer
2 treatment, palliative care, and end-of-life care.” The Guideline also assesses the risks and harms
3 associated with opioid use.

4 326. The 2016 CDC Guideline is the result of a thorough and extensive process by the
5 CDC. The CDC issued the Guideline after it “obtained input from experts, stakeholders, the
6 public, peer reviewers, and a federally chartered advisory committee.” The recommendations in
7 the 2016 CDC Guideline were further made “on the basis of a systematic review of the best
8 available evidence . . .”

9 327. The CDC went through an extensive and detailed process to solicit expert
10 opinions for the Guideline:

11 CDC sought the input of experts to assist in reviewing the evidence and providing
12 perspective on how CDC used the evidence to develop the draft recommendations.
13 These experts, referred to as the “Core Expert Group” (CEG) included subject
14 matter experts, representatives of primary care professional societies and state
15 agencies, and an expert in guideline development methodology. CDC identified
16 subject matter experts with high scientific standing; appropriate academic and
17 clinical training and relevant clinical experience; and proven scientific excellence
18 in opioid prescribing, substance use disorder treatment, and pain management.
19 CDC identified representatives from leading primary care professional
20 organizations to represent the audience for this guideline. Finally, CDC identified
21 state agency officials and representatives based on their experience with state
22 guidelines for opioid prescribing that were developed with multiple agency
23 stakeholders and informed by scientific literature and existing evidence-based
24 guidelines.

25 328. The 2016 Guideline was also peer-reviewed pursuant to “the final information
26 quality bulletin for peer review.” Specifically, the Guideline describes the following independent
peer-review process:

[P]eer review requirements applied to this guideline because it provides influential
scientific information that could have a clear and substantial impact on public- and
private-sector decisions. Three experts independently reviewed the guideline to
determine the reasonableness and strength of recommendations; the clarity with
which scientific uncertainties were clearly identified; and the rationale, importance,
clarity, and ease of implementation of the recommendations. CDC selected peer

1 reviewers based on expertise, diversity of scientific viewpoints, and independence
2 from the guideline development process. CDC assessed and managed potential
3 conflicts of interest using a process similar to the one as described for solicitation
4 of expert opinion. No financial interests were identified in the disclosure and review
process, and nonfinancial activities were determined to be of minimal risk; thus, no
significant conflict of interest concerns were identified.

5 329. The findings in the 2016 CDC Guideline both confirmed the existing body of
6 scientific evidence regarding the questionable efficacy of opioid use and contradicted
7 Defendants' statements about opioids.

8 330. For instance, the Guideline states "[e]xtensive evidence shows the possible harms
9 of opioids (including opioid use disorder, overdose, and motor vehicle injury)" and that "[o]pioid
10 pain medication use presents serious risks, including overdose and opioid use disorder." The
11 Guideline further confirms there are significant symptoms related to opioid withdrawal,
12 including drug cravings, anxiety, insomnia, abdominal pain, vomiting, diarrhea, sweating,
13 tremor, tachycardia (rapid heartbeat), spontaneous abortion and premature labor in pregnant
14 women, and the unmasking of anxiety, depression, and addiction. These findings contradict
15 statements made by Defendants regarding the minimal risks associated with opioid use,
16 including that the risk of addiction from chronic opioid use is low.

17 331. The Guideline also concludes that there is "[n]o evidence" to show "a long-term
18 benefit of opioids in pain and function versus no opioids for chronic pain . . ." Furthermore, the
19 Guideline indicates that "continuing opioid therapy for 3 months substantially increases the risk
20 of opioid use disorder." Indeed, the Guideline indicates that "[p]atients who do not experience
21 clinically meaningful pain relief early in treatment . . . are unlikely to experience pain relief with
22 longer-term use," and that physicians should "reassess[] pain and function within 1 month" in
23 order to decide whether to "minimize risks of long-term opioid use by discontinuing opioids"
24 because the patient is "not receiving a clear benefit." These findings flatly contradict claims
25 made by the Defendants that there are minimal or no adverse effects of long-term opioid use, or
26 that long-term opioid use could actually improve or restore a patient's function.

1 332. In support of these statements about the lack of long-term benefits of opioid use,
2 the CDC concluded that “[a]lthough opioids can reduce pain during short-term use, the clinical
3 evidence review found insufficient evidence to determine whether pain relief is sustained and
4 whether function or quality of life improves with long-term opioid therapy.” The CDC further
5 found that “evidence is limited or insufficient for improved pain or function with long-term use
6 of opioids for several chronic pain conditions for which opioids are commonly prescribed, such
7 as low back pain, headache, and fibromyalgia.”

8 333. With respect to opioid dosing, the Guideline reports that “[b]enefits of high-dose
9 opioids for chronic pain are not established” while the “risks for serious harms related to opioid
10 therapy increase at higher opioid dosage.” The CDC specifically explains that “there is now an
11 established body of scientific evidence showing that overdose risk is increased at higher opioid
12 dosages.” The CDC also states that there is an “increased risk[] for opioid use disorder,
13 respiratory depression, and death at higher dosages.” As a result, the CDC advises doctors to
14 “avoid increasing dosage” above 90 MME per day. These findings contradict statements made
15 by Defendants that increasing dosage is safe and that under-treatment is the cause for certain
16 patients’ aberrant behavior.

17 334. The 2016 CDC Guideline also contradicts statements made by Defendants that
18 there are reliable risk-mitigation tactics to reduce the risk of addiction. For instance, the
19 Guideline indicates that available risk screening tools “show insufficient accuracy for
20 classification of patients as at low or high risk for [opioid] abuse or misuse” and counsels that
21 doctors “should not overestimate the ability of these tools to rule out risks from long-term opioid
22 therapy.”

23 335. Finally, the 2016 CDC Guideline states that “[n]o studies” support the notion that
24 “abuse-deterrent technologies [are] a risk mitigation strategy for deterring or preventing abuse,”
25 noting that the technologies—even when they work—“do not prevent opioid abuse through oral
26

intake, the most common route of opioid abuse, and can still be abused by nonoral routes.” In particular, the CDC found as follows:

The “abuse-deterrent” label does not indicate that there is no risk for abuse. No studies were found in the clinical evidence review assessing the effectiveness of abuse-deterrent technologies as a risk mitigation strategy for deterring or preventing abuse. In addition, abuse-deterrent technologies do not prevent unintentional overdose through oral intake. Experts agreed that recommendations could not be offered at this time related to use of abuse-deterrent formulations.

Accordingly, the CDC’s findings regarding “abuse-deterrent technologies” directly contradict Purdue and Endo’s claims that their new pills deter or prevent abuse.

336. Notably, in addition to the findings made by the CDC in 2016, the Washington State Agency Medical Directors’ Group (AMDG)—a collaboration among several Washington State Agencies—published its *Interagency Guideline on Prescribing Opioids for Pain* in 2015. The AMDG came to many of the same conclusions as the CDC did. For example, the AMDG found that “there is little evidence to support long term efficacy of [chronic opioid analgesic therapy, or “COAT”] in improving function and pain, [but] there is ample evidence of its risk for harm . . .”¹⁹²

337. In addition, as discussed above, in contrast to Defendants’ statements that the 1980 Porter and Jick letter provided evidence of the low risk of opioid addiction in pain patients, the NEJM recently published a letter largely debunking the use of the Porter and Jick letter as evidence for such a claim.¹⁹³ The researchers demonstrated how the Porter and Jick letter was irresponsibly cited and, in some cases, “grossly misrepresented,” when in fact it did not provide evidence supporting the broad claim of low addiction risk for all patients prescribed opioids for pain. As noted above, Dr. Jick reviewed only files of patients administered opioids in a hospital setting, rather than patients sent home with a prescription for opioids to treat chronic pain.

338. The authors of the 2017 letter described their methodology as follows:

¹⁹² *Interagency Guideline on Prescribing Opioids for Pain*, Agency Med. Directors’ Group (June 2015), <http://www.agencymeddirectors.wa.gov/Files/2015AMDGOpioidGuideline.pdf>.

¹⁹³ Leung, et al., *supra* note 110.

We performed a bibliometric analysis of this [1980] correspondence from its publication until March 30, 2017. For each citation, two reviewers independently evaluated the portrayal of the article's conclusions, using an adaptation of an established taxonomy of citation behavior along with other aspects of generalizability . . . For context, we also ascertained the number of citations of other stand-alone letters that were published in nine contemporaneous issues of the *Journal* (in the index issue and in the four issues that preceded and followed it).

We identified 608 citations of the index publication and noted a sizable increase after the introduction of OxyContin (a long-acting formulation of oxycodone) in 1995 . . . **Of the articles that included a reference to the 1980 letter, the authors of 439 (72.2%) cited it as evidence that addiction was rare in patients treated with opioids. Of the 608 articles, the authors of 491 articles (80.8%) did not note that the patients who were described in the letter were hospitalized at the time they received the prescription, whereas some authors grossly misrepresented the conclusions of the letter . . .** Of note, affirmational citations have become much less common in recent years. In contrast to the 1980 correspondence, 11 stand-alone letters that were published contemporaneously by the *Journal* were cited a median of 11 times.¹⁹⁴ (Emphasis added).

339. The researchers provided examples of quotes from articles citing the 1980 letter, and noted several shortcomings and inaccuracies with the quotations. For instance, the researchers concluded that these quotations (i) “overstate[] conclusions of the index publication,” (ii) do[] not accurately specify its study population,” and (iii) did not adequately address “[l]imitizations to generalizability.”¹⁹⁵

¹⁹⁴ *Id.* (emphasis added).

¹⁹⁵ Supplementary Appendix to Pamela T.M. Leung, B.Sc. Pharm., Erin M. Macdonald, M.Sc., Matthew B. Stanbrook, M.D., Ph.D., Irfan Al Dhalla, M.D., David N. Juurlink, M.D., Ph.D., *A 1980 Letter on the Risk of Opioid Addiction*, 376 N Engl J Med 2194-95 (June 1, 2017), http://www.nejm.org/doi/suppl/10.1056/NEJMc1700150/suppl_file/nejmc1700150_appendix.pdf.

Quote	Reference	Comment
"This pain population with no abuse history is literally at no risk for addiction."	Kowal N. What is the issue?: pseudoaddiction or undertreatment of pain. Nurs Econ 1998;17(6):348-9	
"In truth, however, the medical evidence overwhelmingly indicates that properly administered opioid therapy rarely if ever results in "accidental addiction" or "opioid abuse"."	Libby RT. Treating Doctors as Drug Dealers: The Drug Enforcement Administration's War on Prescription Painkillers. The Independent Review 2006;10(4):511-545.	
"Fear of addiction may lead to reluctance by the physician to prescribe. [...] However, there is no evidence that this occurs when prescribing opioids for pain."	Iles S, Catterall JR, Hanks G. Use of opioid analgesics in a patient with chronic abdominal pain. Int J Clin Pract 2002;56(3):227-8.	
"In reality, medical opioid addiction is very rare. In Porter and Jick's study on patients treated with narcotics, only four of the 11,882 cases showed psychological dependency."	Liu W, Xie S, Yue L, et al. Investigation and analysis of oncologists' knowledge of morphine usage in cancer pain treatment. Onco Targets Ther 2014;7:729-37.	Overstates conclusions of the index publication does not accurately specify its study population. Limitations to generalizability are not otherwise explicitly mentioned.
"Physicians are frequently concerned about the potential for addiction when prescribing opiates; however, there have been studies suggesting that addiction rarely evolves in the setting of painful conditions."	Curtis LA, Morrell TD, Todd KH. Pain Management in the Emergency Department 2006;8(7).	
"Although medicine generally regards anecdotal information with disdain (rigorously controlled double-blind clinical trials are the "gold standard"), solid data on the low risk of addiction to opioid analgesics and the manageability of adverse side effects have been ignored or discounted in favor of the anecdotal, the scientifically unsupported, and the clearly fallacious."	Rich BA. Prioritizing pain management in patient care. Has the time come for a new approach. Postgrad Med 2001;110(3):15-7.	
"The Boston Drug Surveillance Program reviewed the charts of nearly 12,000 cancer pain patients treated over a decade and found only four of them could be labeled as addicts."	Levy MH. Pharmacologic management of cancer pain. Semin Oncol 1994;21(6):718-39.	Incorrectly identifies the index study population as cancer patients; does not otherwise address limitations to generalizability.

340. Based on this review, the researchers concluded as follows:

[W]e found that a five-sentence letter published in the Journal in 1980 was heavily and uncritically cited as evidence that addiction was rare with long-term opioid therapy. We believe that this citation pattern contributed to the North American opioid crisis by helping to shape a narrative that allayed prescribers' concerns about the risk of addiction associated with long-term opioid therapy. In 2007, the manufacturer of OxyContin and three senior executives pleaded guilty to federal criminal charges that they misled regulators, doctors, and patients about the risk of addiction associated with the drug. Our findings highlight the potential

1 consequences of inaccurate citation and underscore the need for diligence when
2 citing previously published studies.¹⁹⁶

3 341. These researchers' careful analysis demonstrates the falsity of Defendants' claim
4 that this 1980 letter was evidence of a low risk of addiction in opioid-treated patients. By casting
5 this letter as evidence of low risk of addiction, Defendants played fast and loose with the truth,
6 with blatant disregard for the consequences of their misrepresentations.

7 **G. Jefferson County Has Been Directly Affected by the Opioid Epidemic Caused by**
8 **Defendants.**

9 342. Jefferson County, located in Western Washington State, has over 31,000
10 residents, the majority of whom live in Port Townsend, the county seat.

11 343. Much like the rest of the United States, Jefferson County has felt the profound
12 consequences of the opioid epidemic. As a direct result of Defendants' aggressive marketing
13 scheme and failure to stop the flood of prescription opioids, Jefferson County has suffered
14 significant and ongoing harms—harms that will continue well into the future. Each day that
15 Defendants continue to evade responsibility for the epidemic they caused, the County must
16 continue allocating substantial resources to address it.

17 344. Opioid use has reached crisis levels across the country, and Jefferson County is
18 not immune to national trends. Statistics regarding opioid overdoses and treatment opioid-use
19 disorder—while not capturing the whole story of this far-reaching epidemic—provide
20 quantitative indicators of the extent of the crisis.

21 345. In Jefferson County, the statistics show that, in recent years, the crisis has only
22 grown. For example, between 2002-2004 and 2011-2013, the number of opioid-use treatment
23
24
25

26 ¹⁹⁶ Leung, et al., *supra* note 110.

admissions rose 206.1%.¹⁹⁷ From 1999 to 2003, there were six opioid-related deaths in Jefferson County;¹⁹⁸ this count quadrupled to twenty-four opioid-related deaths between 2008 and 2012.¹⁹⁹

346. In 2014, Jefferson County had an opioid prescribing rate of 96.5 prescriptions per 100 persons.²⁰⁰

347. Another effect of the opioid epidemic in Jefferson County is the increase in the County's homeless population, which has grown in recent years. Although the causes of homelessness are multi-faceted and complex, opioid abuse is both a contributing cause and result of homelessness. Opioid-use disorder is also a significant factor that prevents someone from regaining economic well-being and housing stability.

348. According to the annual Washington State Point-In-Time Count, the number of homeless persons in Washington State grew 18.8% from 2013 to 2017. In Jefferson County in 2013, there were 98 homeless households.²⁰¹ In 2017, that number had nearly doubled to 187 homeless households in the County.²⁰²

349. In the fall of 2016, Jefferson County worked with neighboring Kitsap and Clallam Counties as part of a project known as Olympic Community of Health to initiate the Three-County Coordinated Opioid Response Project (3CCORP), a multi-sector collaborative effort to address the opioid epidemic in the region. Jefferson County provides extensive staff time to support 3CCORP's efforts.

¹⁹⁷ *Opioid Trends Across Washington State*, U. of Wash. Alcohol & Drug Abuse Inst. (Apr. 2015) <http://adaa.uw.edu/pubs/infobriefs/ADAI-IB-2015-01.pdf>

¹⁹⁸ *Washington State Drug Overdose Quarterly Report*, Wash. St. Dep't of Health, <https://www.doh.wa.gov/DataandStatisticalReports/InjuryViolenceandPoisoning/InjuryData/WashingtonStateInjuryDataTables/OpioidQuarterlyReport> (last visited Aug. 13, 2018).

¹⁹⁹ *Id.*

²⁰⁰ *Opioid Overdose, U.S. County Prescribing Rates, 2014*, Ctrs. for Disease Control and Prevention, <https://www.cdc.gov/drugoverdose/maps/rxcounty2014.html> (last updated July 31, 2017).

²⁰¹ *2013 Point in Time Count*, Wash. Dep't of Commerce (Jan. 2013), <http://www.commerce.wa.gov/wp-content/uploads/2016/10/hau-pit-final-summary-2013.pdf>.

²⁰² *2017 Point in Time Count*, Wash. Dep't of Commerce (Aug. 2017), <http://www.commerce.wa.gov/wp-content/uploads/2017/12/CSHD-HAU-2017-County-Summary-August-2017v2.pdf>.

350. Within 3CCORP, members have formed three workgroups to address (1) Overdose Prevention, (2) Treatment, and (3) Prevention. The Overdose Prevention Workgroup's work includes, for example, identifying baseline metrics for overdose response and finding ways to increase training and the availability of naloxone. The Prevention Workgroup is working on reforming prescribing practices through the Six Building Blocks to Safer Opioid Prescribing model,²⁰³ developed by researchers at the University of Washington and Group Health (now Kaiser) Research Institute in Seattle, and is aimed at revising chronic opioid therapy practices in primary care clinics.²⁰⁴

351. The Treatment Workgroup collaborated and successfully obtained Washington State Department of Social and Health Services funding to support a "hub and spoke" model of treatment for opioid-use disorder. Now being adapted and implemented in various regions after its success in Vermont,²⁰⁵ the "hub and spoke" model provides specialized substance-use disorder treatment in a "hub" with care managers in several "spokes." In Washington State's adaptation of the model, the hub must be a primary care clinic, a community-based health center, or an opioid treatment program, capable of providing on site at least two of the three FDA-approved medications for opioid-use disorder (methadone, buprenorphine, and naltrexone), with a minimum of two waived providers, a nurse care manager, and a care coordinator. A minimum of five spokes must each have a care coordinator and offer supportive recovery services such as counseling, but can include locations such as syringe exchange programs or correctional facilities, as the goal of the spokes is offer low barriers for accessing treatment. While the 3CCORP Treatment Workgroup recognized that ideally each of the three counties

²⁰³ *Six Building Blocks*, <https://www.improvingopioidcare.org/> (last visited Aug. 13, 2018).

²⁰⁴ Michael L. Parchman, MD, MPH, Michael Von Korff, PhD, Laura-Mae Baldwin, MD, Mark Stephens, BS, Brooke Ike, MPH, DeAnn Crompt, MPH, Clariss Hsu, PhD, and Ed H. Wagher, MD, MPH, *Primary Care Clinic Re-Design for Prescription Opioid Management*, 30(1) JABFM 44-51 (Jan.-Feb. 2017), https://www.improvingopioidcare.org/wp-content/uploads/2018/02/JABFM-article_2017.pdf.

²⁰⁵ John R. Brooklyn, MD and Stacey C. Sigmon, PhD, *Vermont Hub-and-Spoke Model of Care For Opioid Use Disorder: Development, Implementation, and Impact*, 11(4) J Addict Med 286-292 (July-Aug. 2017), <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC5537005/>.

1 would have a hub, currently available resources limit the model to one hub for the three counties.
2 Peninsula Community Health Services in Kitsap County was selected as the “hub.” As of March
3 2018, the project had already surpassed its treatment induction goal with 260 treatment
4 inductions.

5 352. Jefferson County has been working to confront the emergency caused by
6 Defendants’ reckless promotion and distribution of prescription opioids. The County has spent
7 substantial sums in the past and will continue to spend substantial sums in the future to address
8 the epidemic.

9 **H. No Federal Agency Action, Including by the FDA, Can Provide the Relief Jefferson**
10 **County Seeks Here.**

11 353. The injuries Jefferson County has suffered and will continue to suffer cannot be
12 addressed by agency or regulatory action. There are no rules the FDA could make or actions the
13 agency could take that would provide Jefferson County the relief it seeks in this litigation.

14 354. Even if prescription opioids were entirely banned today or only used for the
15 intended purpose, millions of Americans, including Jefferson County residents, would remain
16 addicted to opioids, and overdoses will continue to claim lives. The County’s public health,
17 criminal justice, and social welfare services will continue to be burdened by the consequences of
18 the epidemic.

19 355. Regulatory action would do nothing to compensate the County for the money and
20 resources it has already expended addressing the impacts of the opioid epidemic and the
21 resources it will need in the future. Only this litigation has the ability to provide the County with
22 the relief it seeks.

23 356. Furthermore, the costs Jefferson County has incurred in responding to the opioid
24 crisis and in rendering public services described above are recoverable pursuant to the causes of
25 actions raised by the County. Defendants’ misconduct alleged herein is not a series of isolated
26 incidents, but instead the result of a sophisticated and complex marketing scheme over the course

1 of more than twenty years that has caused a substantial and long-term burden on the municipal
 2 services provided by the County. In addition, the public nuisance created by Defendants and the
 3 County's requested relief in seeking abatement further compels Defendants to reimburse and
 4 compensate Jefferson County for the substantial resources it has expended to address the opioid
 5 crisis.

6 **V. CLAIMS FOR RELIEF**
 7 **COUNT ONE — VIOLATIONS OF THE WASHINGTON CONSUMER PROTECTION**
 8 **ACT, RCW 19.86, ET SEQ.**

9 357. Plaintiff repeats, reasserts, and incorporates the allegations contained above as if
 10 fully set forth herein.

11 358. The Washington Consumer Protection Act is codified at RCW 19.86 *et seq.*
 12 (CPA). The CPA establishes a comprehensive framework for redressing the violations of
 13 applicable law, and municipalities of Washington State like Jefferson County can enforce the
 14 CPA and recover damages. RCW 19.86.090. The conduct at issue in this case falls within the
 15 scope of the CPA.

16 359. The CPA prohibits unfair methods of competition and unfair or deceptive acts or
 17 practices in the conduct of any trade or commerce. Defendants engaged and continue to engage
 18 in the same pattern of unfair methods of competition, and unfair and/or deceptive conduct
 19 pursuant to a common practice of misleading the public regarding the purported benefits and
 20 risks of opioids.

21 360. Manufacturing Defendants, at all times relevant to this Complaint, directly and/or
 22 through their control of third parties, violated the CPA by making unfair and/or deceptive
 23 representations about the use of opioids to treat chronic and non-cancer pain, including to
 24 physicians and consumers in Jefferson County. Each Manufacturing Defendant also omitted or
 25 concealed material facts and failed to correct prior misrepresentations and omissions about the
 26 purported benefits and risks of opioids. In addition, each Manufacturing Defendant's silence
 regarding the full risks of opioid use constitutes deceptive conduct prohibited by the CPA.

1 361. The Distributor Defendants, at all times relevant to this Complaint, directly and/or
2 through their control of third parties, violated the CPA by making unfair and/or deceptive
3 representations about their compliance with their obligations to maintain effective controls
4 against diversion of prescription opioids and to report suspicious orders. The Distributor
5 Defendants concealed the extent of their opioid distribution in order to avoid the issuance of
6 restrictive quotas, and manipulated the political process to shield themselves from enforcement
7 actions that would have stopped shipments of opioids.

8 362. These unfair methods of competition and unfair and/or deceptive acts or practices
9 in the conduct of trade or commerce were reasonably calculated to deceive Jefferson County and
10 its consumers, and did in fact deceive the County and its consumers. Each Manufacturing
11 Defendant's misrepresentations, concealments, and omissions continue to this day.

12 363. As a result of Defendants' misrepresentations, Jefferson County has spent
13 substantial sums of money on increased law enforcement, emergency services, social services,
14 public safety, and other human services in Jefferson County.

15 364. But for these unfair methods of competition and unfair and/or deceptive acts or
16 practices in the conduct of trade or commerce, Jefferson County would not have incurred the
17 massive costs related to the epidemic caused by Defendants.

18 365. Logic, common sense, justice, policy, and precedent indicate Manufacturing
19 Defendants' unfair and deceptive conduct has caused the damage and harm complained of
20 herein. Manufacturing Defendants knew or reasonably should have known that their statements
21 regarding the risks and benefits of opioids were false and misleading, and that their statements
22 were causing harm from their continued production and marketing of opioids. The Distributor
23 Defendants knew or reasonably should have known that the proliferation of prescription opioids
24 was causing damage to the County. Thus, the harms caused by Defendants' unfair and deceptive
25 conduct to Jefferson County were reasonably foreseeable, including the financial and economic
26 losses incurred by the County.

1 366. Furthermore, Jefferson County brings this cause of action in its sovereign capacity
 2 for the benefit of the State of Washington. The CPA expressly authorizes local governments to
 3 enforce its provisions and to recover damages for violations of the CPA, and this action is
 4 brought to promote the public welfare of the state and for the common good of the state.

5 367. As a direct and proximate cause of each Defendant's unfair and deceptive
 6 conduct, (i) the County has sustained and will continue to sustain injuries, and (ii) pursuant to
 7 RCW 19.86.090, the County is entitled to actual and treble damages in amounts to be determined
 8 at trial, attorneys' fees and costs, and all other relief available under the CPA.

9 368. The Court should also grant injunctive relief enjoining Defendants from future
 10 violations of the CPA. Defendants' actions, as complained of herein, constitute unfair
 11 competition or unfair, deceptive, or fraudulent acts or practices in violation of the CPA.

12 COUNT TWO — PUBLIC NUISANCE

13 369. Plaintiff repeats, reasserts, and incorporates the allegations contained above as if
 14 fully set forth herein.

15 370. Pursuant to RCW 7.48.010, an actionable nuisance is defined as, *inter alia*,
 16 "whatever is injurious to health or indecent or offensive to the senses . . ."

17 371. Pursuant to RCW 7.48.130, "A public nuisance is one which affects equally the
 18 rights of an entire community or neighborhood, although the extent of the damage may be
 19 unequal."

20 372. Plaintiff and its residents have a right to be free from conduct that endangers their
 21 health and safety. Yet Defendants have engaged in conduct which endangers or injures the health
 22 and safety of the residents of the County by their production, promotion, distribution, and
 23 marketing of opioids for use by residents of Jefferson County and in a manner that substantially
 24 interferes with the welfare of Jefferson County.

25 373. Each Defendant has created or assisted in the creation of a condition that is
 26 injurious to the health and safety of Jefferson County and its residents, and interferes with the

1 comfortable enjoyment of life and property of entire communities and/or neighborhoods in the
2 County.

3 374. Defendants' conduct has directly caused deaths, serious injuries, and a severe
4 disruption of the public peace, order and safety, including fueling the homeless and heroin crises
5 facing the County described herein. Defendants' conduct is ongoing and continues to produce
6 permanent and long-lasting damage.

7 375. The health and safety of the residents of Jefferson County, including those who
8 use, have used, or will use opioids, as well as those affected by users of opioids, are matters of
9 substantial public interest and of legitimate concern to the County's citizens and its residents.

10 376. Defendants' conduct has affected and continues to affect a substantial number of
11 people within Jefferson County and is likely to continue causing significant harm to patients with
12 chronic pain who are being prescribed and take opioids, their families, and their communities.

13 377. But for Defendants' actions, opioid use and ultimately its misuse and abuse would
14 not be as widespread as it is today, and the massive epidemic of opioid abuse that currently exists
15 would have been averted.

16 378. Logic, common sense, justice, policy, and precedent indicate Defendants' unfair
17 and deceptive conduct has caused the damage and harm complained of herein. Manufacturing
18 Defendants knew or reasonably should have known that their statements regarding the risks and
19 benefits of opioids were false and misleading, and that their false and misleading statements
20 were causing harm from their continued production and marketing of opioids. Distributor
21 Defendants knew that the widespread distribution of opioids would endanger the health and
22 safety of residents of Jefferson County. Thus, the public nuisance caused by Defendants to
23 Jefferson County was reasonably foreseeable, including the financial and economic losses
24 incurred by the County.

25 379. Furthermore, Jefferson County brings this cause of action in its capacity as a
26 public body authorized by law to prevent, remove, and abate a nuisance at the expense of the

1 parties creating, causing, or committing the nuisance. The applicable RCW with respect to a
2 public nuisance expressly prohibits the conduct complained of herein, and this action is brought
3 to promote the public health of Jefferson County and for the common good of Jefferson County.

4 380. In addition, engaging in any business in defiance of a law regulating or
5 prohibiting the same is a nuisance per se under Washington law. Each Defendant's conduct
6 described herein of deceptively marketing or excessively distributing opioids violates RCW
7 7.48.010 and therefore constitutes a nuisance per se.

8 381. As a direct and proximate cause of Defendants' conduct creating or assisting in
9 the creation of a public nuisance, Jefferson County, its community, and its residents have
10 sustained and will continue to sustain substantial injuries.

11 382. Pursuant to RCW 7.48.020, Plaintiff requests an order providing for abatement of
12 the public nuisance that each Defendant has created or assisted in the creation of, and enjoining
13 Defendants from future violations of RCW 7.48.010

14 383. Plaintiff also seeks the maximum statutory and civil penalties permitted by law as
15 a result of the public nuisance created by Defendants.

16 **COUNT THREE — NEGLIGENCE**

17 384. Plaintiff repeats, reasserts, and incorporates the allegations contained above as if
18 fully set forth herein.

19 385. Under Washington law, a cause of action arises for negligence when a defendant
20 owes a duty to a plaintiff and breaches that duty, and proximately causes the resulting injury.
21 *Iwai v. State*, 129 Wn. 2d 84, 96, 915 P.2d 1089 (1996).

22 386. Each Defendant owed a duty of care to Jefferson County, including but not
23 limited to taking reasonable steps to prevent the misuse, abuse, and over-prescription of opioids.

24 387. In violation of this duty, Defendants failed to take reasonable steps to prevent the
25 misuse, abuse, and over-prescription of opioids in Jefferson County by misrepresenting the risks
26 and benefits associated with opioids and by distributing dangerous quantities of opioids.

1 388. As set forth above, Manufacturing Defendants' misrepresentations include falsely
2 claiming that the risk of opioid addiction was low, falsely instructing doctors and patients that
3 prescribing more opioids was appropriate when patients presented symptoms of addiction,
4 falsely claiming that risk-mitigation strategies could safely address concerns about addiction,
5 falsely claiming that doctors and patients could increase opioid doses indefinitely without added
6 risk, deceptively marketing that purported abuse-deterrent technology could curb misuse and
7 addiction, and falsely claiming that long-term opioid use could actually restore function and
8 improve a patient's quality of life. Each of these misrepresentations made by Defendants violated
9 the duty of care to Jefferson County.

10 389. Distributor Defendants negligently distributed enormous quantities of potent
11 narcotics and failed to report such distributions. Distributor Defendants violated their duty of
12 care by moving these dangerous products into Jefferson County in such quantities, facilitating
13 diversion, misuse, and abuse of opioids.

14 390. As a direct and proximate cause of Defendants' unreasonable and negligent
15 conduct, Plaintiff has suffered and will continue to suffer harm, and is entitled to damages in an
16 amount determined at trial.

17 **COUNT FOUR — GROSS NEGLIGENCE**

18 391. Plaintiff repeats, reasserts, and incorporates the allegations contained above as if
19 fully set forth herein.

20 392. As set forth above, each Defendant owed a duty of care to Jefferson County,
21 including but not limited to taking reasonable steps to prevent the misuse, abuse, and over-
22 prescription of opioids.

23 393. In violation of this duty, each Defendant failed to take reasonable steps to prevent
24 the misuse, abuse, and over-prescription of opioids in Jefferson County by misrepresenting the
25 risks and benefits associated with opioids.

1 394. In addition, each Defendant knew or should have known, and/or recklessly
2 disregarded, that the opioids they manufactured, promoted, and distributed were being used for
3 unintended uses.

4 395. For instance, Defendants failed to exercise slight care to Jefferson County by,
5 *inter alia*, failing to take appropriate action to stop opioids from being used for unintended
6 purposes. Furthermore, despite each Defendant's actual or constructive knowledge of the wide
7 proliferation of prescription opioids in Jefferson County, Defendants took no action to prevent
8 the abuse and diversion of these drugs. In fact, Manufacturing Defendants promoted and actively
9 targeted doctors and their patients through training their sales representatives to encourage
10 doctors to prescribe more opioids.

11 396. Manufacturing Defendants' misrepresentations include falsely claiming that the
12 risk of opioid addiction was low, falsely instructing doctors and patients that prescribing more
13 opioids was appropriate when patients presented symptoms of addiction, falsely claiming that
14 risk-mitigation strategies could safely address concerns about addiction, falsely claiming that
15 doctors and patients could increase opioid doses indefinitely without added risk, deceptively
16 marketing that purported abuse-deterrent technology could curb misuse and addiction, and
17 falsely claiming that long-term opioid use could actually restore function and improve a patient's
18 quality of life. Each of these misrepresentations made by Manufacturing Defendants violated the
19 duty of care to Plaintiff, in a manner that is substantially and appreciably greater than ordinary
20 negligence.

21 397. Distributor Defendants continued to funnel enormous quantities of opioids into
22 Jefferson County, long after they knew that these products were being misused, abused, and
23 diverted. By permitting the movement of such excessive quantities of dangerous narcotics into
24 Jefferson County, Distributor Defendants endangered the health and safety of Jefferson County
25 residents, in a manner that is substantially and appreciably greater than ordinary negligence.
26

1 398. As a direct and proximate cause of each Defendant's gross negligence, Plaintiff
2 has suffered and will continue to suffer harm, and is entitled to damages in an amount
3 determined at trial.

4 **COUNT FIVE — UNJUST ENRICHMENT**

5 399. Plaintiff repeats, reasserts, and incorporates the allegations contained above as if
6 fully set forth herein.

7 400. Each Defendant was required to take reasonable steps to prevent the misuse,
8 abuse, and over-prescription of opioids.

9 401. Rather than prevent or mitigate the wide proliferation of opioids into Jefferson
10 County, each Defendant instead chose to place its monetary interests first and each Defendant
11 profited from prescription opioids sold in Jefferson County.

12 402. Each Defendant also failed to maintain effective controls against the unintended
13 and illegal use of the prescription opioids it manufactured or distributed, again choosing instead
14 to place its monetary interests first.

15 403. Each Defendant therefore received a benefit from the sale and distribution of
16 prescription opioids to and in Jefferson County, and these Defendants have been unjustly
17 enriched at the expense of Plaintiff.

18 404. As a result, Plaintiff is entitled to damages on its unjust enrichment claim in an
19 amount to be proven at trial.

20 **COUNT SIX — VIOLATIONS OF THE RACKETEER INFLUENCED AND CORRUPT**
21 **ORGANIZATIONS ACT ("RICO"), 18 U.S.C. § 1961, ET SEQ.**

22 405. Plaintiff hereby incorporates by reference the allegations contained in the
23 preceding paragraphs of this complaint.

24 406. This claim is brought by Plaintiff against each Defendant for actual damages,
25 treble damages, and equitable relief under 18 U.S.C. § 1964 for violations of 18 U.S.C. § 1961,
26 *et seq.*

1 407. At all relevant times, each Defendant is and has been a “person” within the
2 meaning of 18 U.S.C. § 1961(3), because they are capable of holding, and do hold, “a legal or
3 beneficial interest in property.”

4 408. Plaintiff is a “person,” as that term is defined in 18 U.S.C. § 1961(3), and has
5 standing to sue as it was and is injured in its business and/or property as a result of the
6 Defendants’ wrongful conduct described herein.

7 409. Section 1962(c) makes it “unlawful for any person employed by or associated
8 with any enterprise engaged in, or the activities of which affect, interstate or foreign commerce,
9 to conduct or participate, directly or indirectly, in the conduct of such enterprise’s affairs through
10 a pattern of racketeering activity . . .” 18 U.S.C. § 1962(c).

11 410. Section 1962(d) makes it unlawful for “any person to conspire to violate” Section
12 1962(c), among other provisions. *See* 18 U.S.C. § 1962(d).

13 411. Each Defendant conducted the affairs of an enterprise through a pattern of
14 racketeering activity, in violation of 18 U.S.C. § 1962(c) and § 1962(d).

15 **A. Description of the Defendants’ Enterprises**

16 412. RICO defines an enterprise as “any individual, partnership, corporation,
17 association, or other legal entity, and any union or group of individuals associated in fact
18 although not a legal entity.” 18 U.S.C. § 1961(4).

19 413. Under 18 U.S.C. § 1961(4) a RICO “enterprise” may be an association-in-fact
20 that, although it has no formal legal structure, has (i) a common purpose, (ii) relationships among
21 those associated with the enterprise, and (iii) longevity sufficient to pursue the enterprise’s
22 purpose. *See Boyle v. United States*, 556 U.S. 938, 946 (2009).

23 414. Defendants formed two such association-in-fact enterprises—referred to herein as
24 “the Promotion Enterprise” and “the Diversion Enterprise.”

25 415. The Promotion Enterprise consists of the Manufacturing Defendants, Front
26 Groups, and KOLs. In particular, the Enterprise consists of (a) Defendant Purdue, including its

1 employees and agents, (b) Defendant Endo, including its employees and agents, (c) Defendant
2 Janssen, including its employees and agents, (d) Defendant Cephalon, including its employees
3 and agents, (e) Defendant Actavis, including its employees and agents, and (f) Defendant
4 Mallinckrodt, including its employees and agents (collectively, “Manufacturing Defendants”);
5 certain front groups described above, including but not limited to (a) the American Pain
6 Foundation, including its employees and agents, (b) the American Academy of Pain Medicine,
7 including its employees and agents, and (c) the American Pain Society, including its employees
8 and agents (collectively, the “Front Groups”); and certain Key Opinion Leaders, including but
9 not limited to (a) Dr. Russell Portenoy, (b) Dr. Perry Fine, (c) Dr. Lynn Webster, and (d) Dr.
10 Scott Fishman (collectively, the “KOLs”). The entities in the Promotion Enterprise acted in
11 concert to create demand for prescription opioids.

12 416. Alternatively, each of the above-named Manufacturing Defendants and Front
13 Groups constitutes a single legal entity “enterprise” within the meaning of 18 U.S.C. § 1961(4),
14 through which the members of the enterprise conducted a pattern of racketeering activity. The
15 separate legal status of each member of the Enterprise facilitated the fraudulent scheme and
16 provided a hoped-for shield from liability for Defendants and their co-conspirators.

17 417. Alternatively, each of the Manufacturing Defendants, together with the
18 Distributor Defendants, the Front Groups, and the KOLs, constitute separate, associated-in-fact
19 Enterprises within the meaning of 18 U.S.C. § 1961(4).

20 418. The Diversion Enterprise consists of all Defendants. In particular, the Enterprise
21 consists of (a) Defendant Purdue, including its employees and agents, (b) Defendant Endo,
22 including its employees and agents, (c) Defendant Janssen, including its employees and agents,
23 (d) Defendant Cephalon, including its employees and agents, (e) Defendant Actavis, including its
24 employees and agents, (f) Defendant Mallinckrodt, including its employees and agents, (g)
25 Defendant AmerisourceBergen, including its employees and agents, (h) Defendant Cardinal
26

1 Health, including its employees and agents, and (i) Defendant McKesson, including its
2 employees and agents (collectively, “Defendants”).

3 419. The CSA and its implementing regulations require all manufacturers and
4 distributors of controlled substances, including opioids, to maintain a system to identify and
5 report suspicious orders, including orders of unusual size or frequency, or orders deviating from
6 a normal pattern, and maintain effective controls against diversion of controlled substances. *See*
7 21 U.S.C. § 823; 21 C.F.R. §1301.74(b). The Manufacturing Defendants and the Distributor
8 Defendants alike are required to become “registrants” under the CSA, 21 U.S.C. § 823(a)-(b),
9 and its implementing regulations, which provide that “[e]very person who manufactures,
10 distributes, dispenses, imports, or exports any controlled substance. . . shall obtain a
11 registration[.]” 21 C.F.R. § 1301.11(a). Defendants’ duties as registrants include reporting
12 suspicious orders of controlled substances, which are defined as including “orders of unusual
13 size, orders deviating substantially from a normal pattern, and orders of unusual frequency.” 21
14 C.F.R. § 1301.74(b).

15 420. The Manufacturing Defendants carried out the Diversion Enterprise by
16 incentivizing and supplying suspicious sales of opioids, despite their knowledge that their
17 opioids were being diverted to illicit use, and by failing to notify the DEA of such suspicious
18 orders as required by law. The Distributor Defendants carried out the Diversion Enterprise by
19 failing to maintain effective controls against diversion, intentionally evading their obligation to
20 report suspicious orders to the DEA, and conspiring to prevent limits on the prescription opioids
21 they were oversupplying to communities like Plaintiff.

22 421. The Promotion Enterprise is an ongoing and continuing business organization
23 consisting of “persons” within the meaning of 18 U.S.C. § 1961(3) that created and maintained
24 systematic links for a common purpose: to sell highly addictive opioids for treatment of chronic
25 pain while knowing that opioids have little or no demonstrated efficacy for such pain and have
26 significant risk of addiction, overdose, and death.

1 422. The Distribution Enterprise is an ongoing and continuing business organization
2 consisting of “persons” within the meaning of 18 U.S.C. § 1961(3) that created and maintained
3 systematic links for a common purpose: to distribute highly addictive opioids in quantities that
4 far exceeded amounts that could reasonably be considered medically necessary.

5 423. To accomplish these purposes, the Promotion Enterprise engaged in a
6 sophisticated, well-developed, and fraudulent marketing scheme designed to increase the
7 prescription rate for Defendants’ opioid medications (the “Promotion Scheme”), and the
8 Diversion Enterprise carried out a scheme to systematically disregard, avoid, or frustrate the
9 monitoring and reporting requirements intended to prevent the widespread distribution of
10 dangerous controlled substances (the “Diversion Scheme”). The Promotion Scheme and the
11 Diversion Scheme are collectively referred to as the “Schemes.”

12 **B. The Enterprises Sought to Fraudulently Increase Defendants’ Profits and Revenues**

13 424. At all relevant times, each Defendant was aware of the conduct of the Enterprises,
14 was a knowing and willing participant in that conduct, and reaped profits from that conduct in
15 the form of increased sales and distribution of prescription opioids. In addition, the Front Groups
16 and KOLs received direct payments from the Manufacturing Defendants in exchange for their
17 role in the Promotion Enterprise, and to advance the Promotion Enterprise’s fraudulent
18 marketing scheme.

19 425. The Enterprises engaged in, and their activities affected, interstate and foreign
20 commerce because they involved commercial activities across state boundaries, including but not
21 limited to: (1) the marketing, promotion, and distribution of prescription opioids; (2) advocacy at
22 the state and federal level for change in the law governing the use and prescription of
23 prescription opioids; (3) the issuance of prescriptions and prescription guidelines for opioids; (4)
24 the issuance of fees, bills, and statements demanding payment for prescriptions of opioids; (5)
25 payments, rebates, and chargebacks between Defendants; and (6) the creation of documents,
26

1 reports, and communications related to Defendants' reporting requirements under the CSA and
2 its implementing regulations.

3 426. The persons engaged in the Enterprises are systematically linked through
4 contractual relationships, financial ties, and continuing coordination of activities, as spearheaded
5 by Defendants. With respect to the Promotion Enterprise, each Manufacturing Defendant funded
6 and directed the operations of the KOLs and the Front Groups; in fact, the board of directors of
7 each of the Front Groups are and were full of doctors who were on the Manufacturing
8 Defendants' payrolls, either as consultants or speakers at medical events. Moreover, each
9 Manufacturing Defendant coordinated and, at times, co-funded their activities in furtherance of
10 the goals of the Enterprise. This coordination can also be inferred through the consistent
11 misrepresentations described below. With respect to the Diversion Enterprise, Defendants were
12 financially linked through a system of payments, rebates, and chargebacks.

13 427. In the Promotion Enterprise, there is regular communication between each
14 Manufacturing Defendant, each of the Front Groups, and each KOL in which information
15 regarding the Defendants' scheme to increase opioid prescriptions is shared. Typically, this
16 communication occurred, and continues to occur, through the use of the wires and the mail in
17 which Manufacturing Defendants, the Front Groups, and the KOL share information regarding
18 the operation of the Promotion Enterprise.

19 428. In the Diversion Enterprise, there is regular communication between each
20 Defendant in which information regarding the Defendants' scheme to oversupply opioids and
21 avoid restrictive regulations or quotas is shared. Typically, this communication occurred, and
22 continues to occur, through the use of the wires and the mail in which Defendants share
23 information regarding the operation of the Diversion Enterprise.

24 429. The Enterprises functioned as continuing units for the purposes of executing the
25 Schemes, and when issues arose during the Schemes, each member of the Enterprises agreed to
26 take actions to hide the Schemes and the existence of the Enterprises.

1 430. Each Defendant participated in the operation and management of the Enterprises
2 by directing its affairs as described herein.

3 431. While Defendants participate in, and are members of, the Enterprises, they have
4 an existence separate from the Enterprises, including distinct legal statuses, affairs, offices and
5 roles, officers, directors, employees, and individual personhood.

6 432. Each Manufacturing Defendant orchestrated the affairs of the Promotion
7 Enterprise and exerted substantial control over the Promotion Enterprise by, at least: (1) making
8 misleading statements about the purported benefits, efficacy, and risks of opioids to doctors,
9 patients, the public, and others, in the form of telephonic and electronic communications, CME
10 programs, medical journals, advertisements, and websites; (2) employing sales representatives to
11 promote the use of opioid medications; (3) purchasing and utilizing sophisticated marketing data
12 (e.g., IMS data) to coordinate and refine the Promotion Scheme; (4) employing doctors to serve
13 as speakers at or attend all-expense paid trips to programs emphasizing the benefits of
14 prescribing opioid medications; (5) funding, controlling, and operating the Front Groups,
15 including the American Pain Foundation and the Pain & Policy Studies Group; (6) sponsoring
16 CME programs that claimed that opioid therapy has been shown to reduce pain and depressive
17 symptoms; (7) supporting and sponsoring guidelines indicating that opioid medications are
18 effective and can restore patients' quality of life; (8) retaining KOLs to promote the use of
19 opioids; and (9) concealing the true nature of their relationships with the other members of the
20 Promotion Scheme, and the Promotion Enterprise, including the Front Groups and the KOLs.

21 433. The Front Groups orchestrated the affairs of the Promotion Enterprise and exerted
22 substantial control over the Promotion Enterprise by, at least: (1) making misleading statements
23 about the purported benefits, efficacy, and low risks of opioids described herein; (2) holding
24 themselves out as independent advocacy groups, when in fact their operating budgets are entirely
25 comprised of contributions from opioid drug manufacturers; (3) publishing treatment guidelines
26 that advised the prescription of opioids; (4) sponsoring medical education programs that touted

1 the benefits of opioids to treat chronic pain while minimizing and trivializing their risks; and (5)
2 concealing the true nature of their relationship with the other members of the Promotion
3 Enterprise.

4 434. The KOLs orchestrated the affairs of the Promotion Enterprise and exerted
5 substantial control over the Promotion Enterprise by, at least: (1) making misleading statements
6 about the purported benefits, efficacy, and low risks of opioids; (2) holding themselves out as
7 independent, when in fact they are systematically linked to and funded by opioid drug
8 manufacturers; and (3) concealing the true nature of their relationship with the other members of
9 the Promotion Enterprise.

10 435. Without the willing participation of each member of the Promotion Enterprise, the
11 Promotion Scheme and the Promotion Enterprise's common course of conduct would not have
12 been successful.

13 436. Each Distributor Defendant orchestrated the affairs of the Diversion Enterprise
14 and exerted substantial control over the Diversion Enterprise by, at least: (1) refusing or failing
15 to identify, investigate, or report suspicious orders of opioids to the DEA; (2) providing the
16 Manufacturing Defendants with data regarding their prescription opioid sales, including purchase
17 orders and ship notices; (3) accepting payments from the Manufacturing Defendants in the form
18 of rebates and/or chargebacks; (4) filling suspicious orders for prescription opioids despite
19 having identified them as suspicious and knowing opioids were being diverted into the illicit
20 drug market; (5) working with other members of the Enterprise through groups like the
21 Healthcare Distribution Alliance to ensure the free flow of opioids, including by supporting
22 limits on the DEA's ability to use immediate suspension orders; and (6) concealing the true
23 nature of their relationships with the other members of the Diversion Enterprise.

24 437. Each Manufacturing Defendant orchestrated the affairs of the Diversion
25 Enterprise and exerted substantial control over the Diversion Enterprise by, at least: (1) refusing
26 or failing to identify, investigate, or report suspicious orders of opioids to the DEA; (2) obtaining

1 from the Distributor Defendants data regarding their prescription opioid sales, including
 2 purchase orders and ship notices; (3) providing payments to the Distributor Defendants in the
 3 form of rebates and/or chargebacks; (4) working with other members of the Diversion Enterprise
 4 through groups like the Healthcare Distribution Alliance to ensure the free flow of opioids,
 5 including by supporting limits on the DEA's ability to use immediate suspension orders; and (5)
 6 concealing the true nature of their relationships with the other members of the Diversion
 7 Enterprise.

8 438. Without the willing participation of each member of the Diversion Enterprise, the
 9 Diversion Scheme and the Diversion Enterprise's common course of conduct would not have
 10 been successful.

11 **C. Predicate Acts: Mail and Wire Fraud**

12 439. To carry out, or attempt to carry out, the Schemes, the members of the
 13 Enterprises, each of whom is a person associated-in-fact with the Enterprises, did knowingly
 14 conduct or participate in, directly or indirectly, the affairs of the Enterprises through a pattern of
 15 racketeering activity within the meaning of 18 U.S.C. §§ 1961(1), 1961(5) and 1962(c), and
 16 employed the use of the mail and wire facilities, in violation of 18 U.S.C. § 1341 (mail fraud)
 17 and § 1343 (wire fraud).

18 440. Specifically, the members of the Enterprises have committed, conspired to
 19 commit, and/or aided and abetted in the commission of, at least two predicate acts of
 20 racketeering activity (i.e., violations of 18 U.S.C. §§ 1341 and 1343), within the past ten years.

21 441. The multiple acts of racketeering activity which the members of the Enterprises
 22 committed, or aided or abetted in the commission of, were related to each other, posed a threat of
 23 continued racketeering activity, and therefore constitute a "pattern of racketeering activity."

24 442. The racketeering activity was made possible by the Enterprises' regular use of the
 25 facilities, services, distribution channels, and employees of the Enterprises.
 26

1 443. The members of the Enterprises participated in the Schemes by using mail,
2 telephone, and the internet to transmit mailings and wires in interstate or foreign commerce.

3 444. The members of the Enterprises used, directed the use of, and/or caused to be
4 used, thousands of interstate mail and wire communications in service of their Schemes through
5 common misrepresentations, concealments, and material omissions.

6 445. In devising and executing the illegal Schemes, the members of the Enterprises
7 devised and knowingly carried out a material scheme and/or artifice to defraud Plaintiff and the
8 public to obtain money by means of materially false or fraudulent pretenses, representations,
9 promises, or omissions of material facts.

10 446. For the purpose of executing the illegal Schemes, the members of the Enterprises
11 committed these racketeering acts, which number in the thousands, intentionally and knowingly
12 with the specific intent to advance the illegal Schemes.

13 447. The Enterprises' predicate acts of racketeering (18 U.S.C. § 1961(1)) include, but
14 are not limited to:

15 A. Mail Fraud: The members of the Enterprises violated 18 U.S.C. § 1341 by
16 sending or receiving, or by causing to be sent and/or received, fraudulent materials
17 via U.S. mail or commercial interstate carriers for the purpose of selling and
18 distributing excessive quantities of highly addictive opioids.

19 B. Wire Fraud: The members of the Enterprises violated 18 U.S.C. § 1343 by
20 transmitting and/or receiving, or by causing to be transmitted and/or received,
21 fraudulent materials by wire for the purpose of selling and distributing excessive
22 quantities of highly addictive opioids.

23 448. The Manufacturing Defendants falsely and misleadingly used the mails and wires
24 in violation of 18 U.S.C. § 1341 and § 1343. Illustrative and non-exhaustive examples include
25 the following: Defendant Purdue's (1) May 31, 1996 press release announcing the release of
26 OxyContin and indicating that the fear of OxyContin's addictive properties was exaggerated; (2)
1990 promotional video in which Dr. Portenoy, a paid Purdue KOL, understated the risk of
opioid addiction; (3) 1998 promotional video which misleadingly cited a 1980 NEJM letter in

1 support of the use of opioids to treat chronic pain; (4) statements made on its 2000 “Partners
2 Against Pain” website which claimed that the addiction risk of OxyContin was very low; (5)
3 literature distributed to physicians which misleadingly cited a 1980 NEJM letter in support of the
4 use of opioids to treat chronic pain; (6) August 2001 statements to Congress by Purdue
5 Executive Vice President and Chief Operating Officer Michael Friedman regarding the value of
6 OxyContin in treating chronic pain; (7) patient brochure entitled “A Guide to Your New Pain
7 Medicine and How to Become a Partner Against Pain” indicating that OxyContin is non-
8 addicting; (8) 2001 statement by Senior Medical Director for Purdue, Dr. David Haddox,
9 indicating that the ‘legitimate’ use of OxyContin would not result in addiction; (9) multiple sales
10 representatives’ communications regarding the low risk of addiction associated with opioids;
11 (10) statements included in promotional materials for opioids distributed to doctors via the mail
12 and wires; (11) statements in a 2003 Patient Information Guide distributed by Purdue indicating
13 that addiction to opioid analgesics in properly managed patients with pain has been reported to
14 be rare; (12) telephonic and electronic communications to doctors and patients indicating that
15 signs of addiction in the case of opioid use are likely only the signs of under-treated pain; (13)
16 statements in Purdue’s Risk Evaluation and Mitigation Strategy for OxyContin indicating that
17 drug-seeking behavior on the part of opioid patients may, in fact, be pain-relief seeking behavior;
18 (14) statements made on Purdue’s website and in a 2010 “Dear Healthcare Professional” letter
19 indicating that opioid dependence can be addressed by dosing methods such as tapering; (15)
20 statements included in a 1996 sales strategy memo indicating that there is no ceiling dose for
21 opioids for chronic pain; (16) statements on its website that abuse-resistant products can prevent
22 opioid addiction; (17) statements made in a 2012 series of advertisements for OxyContin
23 indicating that long-term opioid use improves patients’ function and quality of life; (18)
24 statements made in advertising and a 2007 book indicating that pain relief from opioids improve
25 patients’ function and quality of life; (19) telephonic and electronic communications by its sales
26 representatives indicating that opioids will improve patients’ function; and (20) electronic and

1 telephonic communications concealing its relationship with the other members of the
2 Enterprises.

3 449. Defendant Endo Pharmaceuticals, Inc. also made false or misleading claims in
4 violation of 18 U.S.C. § 1341 and § 1343 including but not limited to: (1) statements made,
5 beginning in at least 2009, on an Endo-sponsored website, PainKnowledge.com, indicating that
6 patients who take opioids as prescribed usually do not become addicted; (2) statements made on
7 another Endo-sponsored website, PainAction.com, indicating that most chronic pain patients do
8 not become addicted to opioid medications; (3) statements in pamphlets and publications
9 described by Endo indicating that most people who take opioids for pain relief do not develop an
10 addiction; (4) statements made on the Endo-run website, Opana.com, indicating that opioid use
11 does not result in addiction; (5) statements made on the Endo-run website, Opana.com,
12 indicating that opioid dependence can be addressed by dosing methods such as tapering; (6)
13 statements made on its website, PainKnowledge.com, that opioid dosages could be increased
14 indefinitely; (7) statements made in a publication entitled “Understanding Your Pain: Taking
15 Oral Opioid Analgesics” suggesting that opioid doses can be increased indefinitely; (8)
16 electronic and telephonic communications to its sales representatives indicating that the formula
17 for its medicines is ‘crush resistant;’ (9) statements made in advertisements and a 2007 book
18 indicating that pain relief from opioids improves patients’ function and quality of life; (10)
19 telephonic and electronic communications by its sales representatives indicating that opioids will
20 improve patients’ function; and (11) telephonic and electronic communications concealing its
21 relationship with the other members of the Enterprises.

22 450. Defendant Janssen made false or misleading claims in violation of 18 U.S.C. §
23 1341 and § 1343 including but not limited to: (1) statements on its website,
24 PrescribeResponsibly.com, indicating that concerns about opioid addiction are overestimated; (2)
25 statements in a 2009 patient education guide claiming that opioids are rarely addictive when used
26 properly; (3) statements included on a 2009 Janssen-sponsored website promoting the concept of

1 opioid pseudoaddiction; (4) statements on its website, PrescribeResponsibly.com, advocating the
2 concept of opioid pseudoaddiction; (5) statements on its website, PrescribeResponsibly.com,
3 indicating that opioid addiction can be managed; (6) statements in its 2009 patient education
4 guide indicating the risks associated with limiting the dosages of pain medicines; (7) telephonic
5 and electronic communications by its sales representatives indicating that opioids will improve
6 patients' function; and (8) telephonic and electronic communications concealing its relationship
7 with the other members of the Enterprises.

8 451. The American Academic of Pain Medicine made false or misleading claims in
9 violation of 18 U.S.C. § 1341 and § 1343 including but not limited to: (1) statements made in a
10 2009 patient education video entitled "Finding Relief: Pain Management for Older Adults"
11 indicating the opioids are rarely addictive; and (2) telephonic and electronic communications
12 concealing its relationship with the other members of the Promotion Enterprise.

13 452. The American Pain Society Quality of Care Committee made a number of false or
14 misleading claims in violation of 18 U.S.C. § 1341 and § 1343 including but not limited to: (1) a
15 May 31, 1996 press release in which the organization claimed there is very little risk of addiction
16 from the proper use of drugs for pain relief; and (2) telephonic and electronic communications
17 concealing its relationship with the other members of the Promotion Enterprise.

18 453. The American Pain Foundation ("APF") made a number of false and misleading
19 claims in violation of 18 U.S.C. § 1341 and § 1343 including but not limited to: (1) statements
20 made by an APF Executive Director to Congress indicating that opioids only rarely lead to
21 addiction; (2) statements made in a 2002 amicus curiae brief filed with an Ohio appeals court
22 claiming that the risk of abuse does not justify restricting opioid prescriptions for the treatment
23 of chronic pain; (3) statements made in a 2007 publication entitled "Treatment Options: A Guide
24 for People Living with Pain" indicating that the risks of addiction associated with opioid
25 prescriptions have been overstated; (4) statements made in a 2002 court filing indicating that
26 opioid users are not "actual addicts"; (5) statements made in a 2007 publication entitled

1 “Treatment Options: A Guide for People Living with Pain” indicating that even physical
2 dependence on opioids does not constitute addiction; (6) claims on its website that there is no
3 ceiling dose for opioids for chronic pain; (7) statements included in a 2011 guide indicating that
4 opioids can improve daily function; and (8) telephonic and electronic communications
5 concealing its relationship with the other members of the Promotion Enterprise.

6 454. The KOLs, including Drs. Russell Portenoy, Perry Fine, Scott Fishman, and Lynn
7 Webster, made a number of misleading statements in the mail and wires in violation of 18 U.S.C.
8 § 1341 and § 1343, described above, including statements made by Dr. Portenoy in a
9 promotional video indicating that the likelihood of addiction to opioid medications is extremely
10 low. Indeed, Dr. Portenoy has since admitted that his statements about the safety and efficacy of
11 opioids were false.

12 455. The Manufacturing Defendants and Distributor Defendants falsely and
13 misleadingly used the mails and wires in violation of 18 U.S.C. § 1341 and § 1343. Illustrative
14 and non-exhaustive examples include the following: (1) the transmission of documents and
15 communications regarding the sale, shipment, and delivery of excessive quantities of
16 prescription opioids, including invoices and shipping records; (2) the transmission of documents
17 and communications regarding their requests for higher aggregate production quotas, individual
18 manufacturing quotas, and procurement quotas; (3) the transmission of reports to the DEA that
19 did not disclose suspicious orders as required by law; (4) the transmission of documents and
20 communications regarding payments, rebates, and chargebacks; (5) the transmission of the actual
21 payments, rebates, and chargebacks themselves; (6) correspondence between Defendants and
22 their representatives in front groups and trade organizations regarding efforts to curtail
23 restrictions on opioids and hobble DEA enforcement actions; (7) the submission of false and
24 misleading certifications required annually under various agreements between Defendants and
25 federal regulators; and (8) the shipment of vast quantities of highly addictive opioids. Defendants
26 also communicated by U.S. mail, by interstate facsimile, and by interstate electronic mail and

1 with various other affiliates, regional offices, regulators, distributors, and other third-party
2 entities in furtherance of the scheme.

3 456. In addition, the Distributor Defendants misrepresented their compliance with laws
4 requiring them to identify, investigate, and report suspicious orders of prescription opioids and/or
5 diversion into the illicit market. At the same time, the Distributor Defendants misrepresented the
6 effectiveness of their monitoring programs, their ability to detect suspicious orders, their
7 commitment to preventing diversion of prescription opioids, and their compliance with
8 regulations regarding the identification and reporting of suspicious orders of prescription opioids.

9 457. The mail and wire transmissions described herein were made in furtherance of
10 Defendants' Schemes and common course of conduct designed to sell drugs that have little or no
11 demonstrated efficacy for the pain they are purported to treat in the majority of persons
12 prescribed them; increase the prescription rate for opioid medications; and popularize the
13 misunderstanding that the risk of addiction to prescription opioids is low when used to treat
14 chronic pain, and to deceive regulators and the public regarding Defendants' compliance with
15 their obligations to identify and report suspicious orders of prescription opioids, while
16 Defendants intentionally enabled millions of prescription opioids to be deposited into
17 communities across the United States, including in Jefferson County. Defendants' scheme and
18 common course of conduct was intended to increase or maintain high quotas for the manufacture
19 and distribution of prescription opioids and their corresponding high profits for all Defendants.

20 458. Many of the precise dates of the fraudulent uses of the U.S. mail and interstate
21 wire facilities have been deliberately hidden, and cannot be alleged without access to
22 Defendants' books and records. However, Plaintiff has described the types of predicate acts of
23 mail and/or wire fraud, including certain specific fraudulent statements and specific dates upon
24 which, through the mail and wires, Defendants engaged in fraudulent activity in furtherance of
25 the Schemes.

1 459. The members of the Enterprises have not undertaken the practices described
2 herein in isolation, but as part of a common scheme and conspiracy. In violation of 18 U.S.C. §
3 1962(d), the members of the Enterprises conspired to violate 18 U.S.C. § 1962(c), as described
4 herein. Various other persons, firms, and corporations, including third-party entities and
5 individuals not named as defendants in this Complaint, have participated as co-conspirators with
6 Defendants and the members of the Enterprises in these offenses and have performed acts in
7 furtherance of the conspiracy to increase or maintain revenue, increase market share, and/or
8 minimize losses for the Defendants and their named and unnamed co-conspirators throughout the
9 illegal scheme and common course of conduct.

10 460. The members of the Enterprises aided and abetted others in the violations of the
11 above laws.

12 461. To achieve their common goals, the members of the Enterprises hid from Plaintiff
13 and the public: (1) the fraudulent nature of the Manufacturing Defendants' marketing scheme;
14 (2) the fraudulent nature of statements made by Defendants and on behalf of Defendants
15 regarding the efficacy of and risk of addiction associated with prescription opioids; (3) the
16 fraudulent nature of the Distributor Defendants' representations regarding their compliance with
17 requirements to maintain effective controls against diversion and report suspicious orders of
18 opioids; and (4) the true nature of the relationship between the members of the Enterprises.

19 462. Defendants and each member of the Enterprises, with knowledge and intent,
20 agreed to the overall objectives of the Schemes and participated in the common course of
21 conduct. Indeed, for the conspiracy to succeed, each of the members of the Enterprises and their
22 co-conspirators had to agree to conceal their fraudulent scheme.

23 463. The members of the Enterprises knew, and intended that, Plaintiff and the public
24 would rely on the material misrepresentations and omissions made by them and suffer damages
25 as a result.
26

1 464. As described herein, the members of the Enterprises engaged in a pattern of
2 related and continuous predicate acts for years. The predicate acts constituted a variety of
3 unlawful activities, each conducted with the common purpose of obtaining significant monies
4 and revenues from Plaintiff and the public based on their misrepresentations and omissions.

5 465. The predicate acts also had the same or similar results, participants, victims, and
6 methods of commission.

7 466. The predicate acts were related and not isolated events.

8 467. The true purposes of Defendants' Schemes were necessarily revealed to each
9 member of the Enterprises. Nevertheless, the members of the Enterprises continued to
10 disseminate misrepresentations regarding the nature of prescription opioids and the functioning
11 of the Schemes.

12 468. Defendants' fraudulent concealment was material to Plaintiff and the public. Had
13 the members of the Enterprises disclosed the true nature of prescription opioids and their
14 excessive distribution, Jefferson County would not have acted as it did or incurred the substantial
15 costs in responding to the crisis caused by Defendants' conduct.

16 469. The pattern of racketeering activity described above is currently ongoing and
17 open-ended, and threatens to continue indefinitely unless this Court enjoins the racketeering
18 activity.

19 **D. Jefferson County Has Been Damaged by Defendants' RICO Violations**

20 470. By reason of, and as a result of the conduct of the Enterprises and, in particular,
21 their patterns of racketeering activity, Jefferson County has been injured in its business and/or
22 property in multiple ways, including but not limited to increased health care costs, increased
23 human services costs, costs related to dealing with opioid-related crimes and emergencies, and
24 other public safety costs, as fully described above.

25 471. Defendants' violations of 18 U.S.C. § 1962(c) and (d) have directly and
26 proximately caused injuries and damages to Jefferson County, its community, and the public, and

the County is entitled to bring this action for three times its actual damages, as well as injunctive/equitable relief, costs, and reasonable attorney's fees pursuant to 18 U.S.C. § 1964(c).

PRAYER FOR RELIEF

WHEREFORE, Plaintiff Jefferson County respectfully requests the Court order the following relief:

- A. An Order that the conduct alleged herein violates the Washington CPA;
- B. An Order that Plaintiff is entitled to treble damages pursuant to the Washington CPA;
- C. An Order that the conduct alleged herein constitutes a public nuisance, including under RCW 7.48 *et seq.*, and under Washington law;
- D. An Order that Defendants abate the public nuisance that they caused;
- E. An Order that Defendants are liable for civil and statutory penalties to the fullest extent permissible under Washington law for the public nuisance they caused;
- F. An Order that Defendants are negligent under Washington law;
- G. An Order that Defendants are grossly negligent under Washington law;
- H. An Order that Defendants have been unjustly enriched at Plaintiff's expense under Washington law;
- I. An Order that Defendants' conduct constitutes violations of the Racketeer Influenced and Corrupt Organizations Act ("RICO"), 18 U.S.C. §1961, *et seq.*;
- J. An Order that Plaintiff is entitled to recover all measure of damages permissible under the statutes identified herein and under common law;
- K. An Order that Defendants are enjoined from the practices described herein;
- L. An Order that judgment be entered against Defendants in favor of Plaintiff;
- M. An Order that Plaintiff is entitled to attorneys' fees and costs pursuant to any applicable provision of law, including but not limited to under the Washington CPA; and

N. An Order awarding any other and further relief deemed just and proper, including pre-judgment and post-judgment interest on the above amounts.

JURY TRIAL DEMAND

Plaintiff demands a trial by jury on all claims and of all issues so triable.

DATED this 14th day of August, 2018.

JEFFERSON COUNTY

KELLER ROHRBACK L.L.P.

By /s/ Philip C. Hunsucker
Philip C. Hunsucker, WSBA #48692
Chief Civil Deputy Prosecuting Attorney
Jefferson County Prosecuting Attorney's Office
P.O. Box 1220
Port Townsend, WA 98368
Phone: (360) 385-9180
Fax: (360) 385-0073

By /s/ Lynn Lincoln Sarko
By /s/ Derek W. Loeser
By /s/ Gretchen Freeman Cappio
By /s/ David J. Ko
By /s/ Daniel P. Mensher
By /s/ Alison S. Gaffney
By /s/ Erika M. Keech
Lynn Lincoln Sarko, WSBA #16569
Derek W. Loeser, WSBA #24274
Gretchen Freeman Cappio, WSBA #29576
David J. Ko, WSBA #38299
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Attorneys for Plaintiff

CIVIL COVER SHEET

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

I. (a) PLAINTIFFS

Jefferson County

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

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

Keller Rohrbach L.L.P.
1201 Third Ave., Suite 3200
Seattle, WA 98101
(206) 623-1900

Jefferson County Prosecuting Attorney's Office
P.O. Box 1220
Port Townsend, WA 98368
(360) 385-9180

DEFENDANTS

Purdue Pharma, L.P.; Purdue Pharma, Inc.; The Purdue Frederick Company, Inc.; Endo Health Solutions Inc.; Endo Pharmaceuticals, Inc.; Janssen Pharmaceuticals, Inc.; Johnson & Johnson; Teva Pharmaceuticals Industries, Ltd.; Teva Pharmaceuticals USA, Inc.; Cephalon, Inc.; Allergan Plc. F/K/A Actavis PLC; Allergan Finance, LLC F/K/A Actavis, Inc. F/K/A Watson Pharmaceuticals, Inc.; Watson Laboratories, Inc.; Actavis LLC; Actavis Pharma, Inc. F/K/A Watson Pharma, Inc.; Mallinckrodt, PLC; Mallinckrodt, LLC; Cardinal Health, Inc.; McKesson Corporation; Amerisourcebergen Drug Corporation; and John and Jane Does 1 through 100, Inclusive

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)

- | | PTF | DEF | | PTF | DEF |
|---|----------------------------|----------------------------|---|----------------------------|----------------------------|
| Citizen of This State | ☐ 1 | ☐ 1 | Incorporated or Principal Place of Business In This State | ☐ 4 | ☐ 4 |
| Citizen of Another State | ☐ 2 | ☐ 2 | Incorporated and Principal Place of Business In Another State | ☐ 5 | ☐ 5 |
| Citizen or Subject of a Foreign Country | ☐ 3 | ☐ 3 | Foreign Nation | ☐ 6 | ☐ 6 |

IV. NATURE OF SUIT (Place an "X" in One Box Only)Click here for: [Nature of Suit Code Descriptions.](#)

CONTRACT	TORTS	FORFEITURE/PENALTY	BANKRUPTCY	OTHER STATUTES
☐ 110 Insurance ☐ 120 Marine ☐ 130 Miller Act ☐ 140 Negotiable Instrument ☐ 150 Recovery of Overpayment & Enforcement of Judgment ☐ 151 Medicare Act ☐ 152 Recovery of Defaulted Student Loans (Excludes Veterans) ☐ 153 Recovery of Overpayment of Veteran's Benefits ☐ 160 Stockholders' Suits ☐ 190 Other Contract ☐ 195 Contract Product Liability ☐ 196 Franchise	PERSONAL INJURY ☐ 310 Airplane ☐ 315 Airplane Product Liability ☐ 320 Assault, Libel & Slander ☐ 330 Federal Employers' Liability ☐ 340 Marine ☐ 345 Marine Product Liability ☐ 350 Motor Vehicle ☐ 355 Motor Vehicle Product Liability ☐ 360 Other Personal Injury ☐ 362 Personal Injury - Medical Malpractice PERSONAL INJURY ☐ 365 Personal Injury - Product Liability ☐ 367 Health Care/Pharmaceutical Personal Injury Product Liability ☐ 368 Asbestos Personal Injury Product Liability PERSONAL PROPERTY ☐ 370 Other Fraud ☐ 371 Truth in Lending ☐ 380 Other Personal Property Damage ☐ 385 Property Damage Product Liability	☐ 625 Drug Related Seizure of Property 21 USC 881 ☐ 690 Other LABOR ☐ 710 Fair Labor Standards Act ☐ 720 Labor/Management Relations ☐ 740 Railway Labor Act ☐ 751 Family and Medical Leave Act ☐ 790 Other Labor Litigation ☐ 791 Employee Retirement Income Security Act IMMIGRATION ☐ 462 Naturalization Application ☐ 465 Other Immigration Actions	☐ 422 Appeal 28 USC 158 ☐ 423 Withdrawal 28 USC 157 PROPERTY RIGHTS ☐ 820 Copyrights ☐ 830 Patent ☐ 835 Patent - Abbreviated New Drug Application ☐ 840 Trademark SOCIAL SECURITY ☐ 861 HIA (1395ff) ☐ 862 Black Lung (923) ☐ 863 DIWC/DIWW (405(g)) ☐ 864 SSID Title XVI ☐ 865 RSI (405(g)) FEDERAL TAX SUITS ☐ 870 Taxes (U.S. Plaintiff or Defendant) ☐ 871 IRS—Third Party 26 USC 7609	☐ 375 False Claims Act ☐ 376 Qui Tam (31 USC 3729(a)) ☐ 400 State Reapportionment ☐ 410 Antitrust ☐ 430 Banks and Banking ☐ 450 Commerce ☐ 460 Deportation ☒ 470 Racketeer Influenced and Corrupt Organizations ☐ 480 Consumer Credit ☐ 490 Cable/Sat TV ☐ 850 Securities/Commodities/Exchange ☐ 890 Other Statutory Actions ☐ 891 Agricultural Acts ☐ 893 Environmental Matters ☐ 895 Freedom of Information Act ☐ 896 Arbitration ☐ 899 Administrative Procedure Act/Review or Appeal of Agency Decision ☐ 950 Constitutionality of State Statutes
REAL PROPERTY ☐ 210 Land Condemnation ☐ 220 Foreclosure ☐ 230 Rent Lease & Ejectment ☐ 240 Torts to Land ☐ 245 Tort Product Liability ☐ 290 All Other Real Property	CIVIL RIGHTS ☐ 440 Other Civil Rights ☐ 441 Voting ☐ 442 Employment ☐ 443 Housing/Accommodations ☐ 445 Amer. w/Disabilities - Employment ☐ 446 Amer. w/Disabilities - Other ☐ 448 Education PRISONER PETITIONS Habeas Corpus: ☐ 463 Alien Detainee ☐ 510 Motions to Vacate Sentence ☐ 530 General ☐ 535 Death Penalty Other: ☐ 540 Mandamus & Other ☐ 550 Civil Rights ☐ 555 Prison Condition ☐ 560 Civil Detainee - Conditions of Confinement			

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):

18 U.S.C. Section 1961, et seq.

Brief description of cause:

Plaintiff alleges violations of the Washington CPA and RICO, and negligence and public nuisance under WA law

VII. REQUESTED IN COMPLAINT:

☐ CHECK IF THIS IS A CLASS ACTION UNDER RULE 23, F.R.Cv.P. DEMAND \$

CHECK YES only if demanded in complaint:

JURY DEMAND: ☒ Yes ☐ No**VIII. RELATED CASE(S)**

IF ANY

(See instructions):

JUDGE Dan A. Polster N.D. OhioDOCKET NUMBER 1:17-md-02804-DAP

DATE

08/14/2018

SIGNATURE OF ATTORNEY OF RECORD

s/ Derek W. Loeser

FOR OFFICE USE ONLY

RECEIPT #

AMOUNT

APPLYING IFP

JUDGE

MAG. JUDGE

UNITED STATES DISTRICT COURT

for the

Western District of Washington

Jefferson County

Plaintiff(s)

v.

Purdue Pharma, L.P.; Purdue Pharma, Inc.; The Purdue Frederick Company, Inc.; Endo Health Solutions Inc.; Endo Pharmaceuticals, Inc.; Janssen Pharmaceuticals, Inc.; Johnson & Johnson; Teva Pharmaceuticals Industries, Ltd.; Teva Pharmaceuticals USA, Inc.; Cephalon, Inc.; Allergan PLC F/K/A Actavis PLC; Allergan Finance, LLC F/K/A Actavis, Inc. F/K/A Watson Pharmaceuticals, Inc.; Watson Laboratories, Inc.; Actavis LLC; Actavis Pharma, Inc. F/K/A Watson Pharma, Inc.; Mallinckrodt, PLC; Mallinckrodt, LLC; Cardinal Health, Inc.; McKesson Corporation; Amerisourcebergen Drug Corporation; and John and Jane Does 1 through 100, Inclusive

Defendant(s)

Civil Action No. 3:18-cv-05661

SUMMONS IN A CIVIL ACTION

To: *(Defendant's name and address)* Purdue Pharma, L.P.; Purdue Pharma, Inc.; The Purdue Frederick Company, Inc.; Endo Health Solutions Inc.; Endo Pharmaceuticals, Inc.; Janssen Pharmaceuticals, Inc.; Johnson & Johnson; Teva Pharmaceuticals Industries, Ltd.; Teva Pharmaceuticals USA, Inc.; Cephalon, Inc.; Allergan PLC F/K/A Actavis PLC; Allergan Finance, LLC F/K/A Actavis, Inc. F/K/A Watson Pharmaceuticals, Inc.; Watson Laboratories, Inc.; Actavis LLC; Actavis Pharma, Inc. F/K/A Watson Pharma, Inc.; Mallinckrodt, PLC; Mallinckrodt, LLC; Cardinal Health, Inc.; McKesson Corporation; Amerisourcebergen Drug Corporation; and John and Jane Does 1 through 100, Inclusive

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:

Derek Loeser
Keller Rohrback L.L.P.
1201 Third Avenue, Suite 3200
Seattle, WA 98101

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. 3:18-cv-05661

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 \$ 0.00 .

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: