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9	IN THE UNITED STAT	ES DISTRICT COURT
10	FOR THE NORTHERN DIS	STRICT OF CALIFORNIA
11	SAN FRANCIS	CO DIVISION
12		
13	STATE OF ALABAMA STATE OF ARKANSAS	SETTLEMENT AGREEMENT AND STIPULATED ORDER FOR
14	STATE OF FLORIDA STATE OF HAWAII	PERMANENT INJUNCTION AND
15	STATE OF IDAHO STATE OF IDAHO STATE OF INDIANA	MONETARY RELIEF
16	STATE OF ILLINOIS	
17	STATE OF IOWA STATE OF MARYLAND	Case No. 3:19-cv-4157
18	STATE OF MINNESOTA STATE OF MISSISSIPPI	
19	STATE OF MISSOURI STATE OF OKLAHOMA	
20	STATE OF OHIO STATE OF UTAH	
21	STATE OF WASHINGTON STATE OF WISCONSIN	
22	COMMONWEALTH OF VIRGINIA	
23	Plaintiffs,	
24	v. ENDO INTERNATIONAL PLC, and	
25	ENDO PHARMACEUTICALS INC.,	
26	Defendants.	
27		
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	1	
	Joint Motion for Entry of Stipulate	d Order for Permanent Injunction

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1	This Settlement Agreement and Stipulated Order ("ASO") is made and entered into this
2	15 th day of July, 2019 ("Effective Date"), by and between the undersigned States through their
3	respective Attorneys General ("Party States"), and Endo Pharmaceuticals Inc. and Endo
4	International plc (collectively "Endo") (the Party States and Endo, collectively, the "Parties").
5	Any State electing to join the ASO shall do so by executing a signature page that shall be annexed
6	to this ASO, with the ASO and all such counterparts collectively constituting the whole of this
7	agreement.
8	WHEREAS, Endo Pharmaceuticals Inc. is or was engaged in manufacturing, marketing,
9	and/or selling certain pharmaceuticals, including but not limited to Lidoderm®, throughout the
10	United States;
11	WHEREAS, the Party States allege that they have reason to believe that Endo
12	Pharmaceuticals Inc. entered into an agreement that foreclosed competition from generic
13	equivalents of the brand-name drug Lidoderm [®] for a period of time in violation of federal
14	antitrust and state antitrust and consumer protection laws;
15	WHEREAS, the Party States initiated an investigation of Endo with respect to the above
16	alleged actions;
17	WHEREAS, the Party States have filed an enforcement Complaint ("Complaint")
18	against Endo in this Court alleging such violations of federal and state laws;
19	WHEREAS, Endo denies that it engaged in any wrongful or unlawful conduct and asserts
20	that it has, at all times, operated within the law and within industry standard practices; and
21	WHEREAS, nothing in this ASO will be construed as a finding or admission of any
22	violation of law on the part of Endo;
23	The Parties have agreed to resolve the States' concerns identified through the
24	investigation and the allegations in the Complaint by entering into this ASO, as follows:
25	FINDINGS
26	1. This Court has jurisdiction over the Parties and the subject matter of this action.
27	Endo has stipulated that, for purposes of this ASO alone, the Court has jurisdiction over Endo
28	Pharmaceuticals Inc. and Endo International plc.
	2 Joint Motion for Entry of Stipulated Order for Permanent Injunction
	Joint Motion for Entry of Supulated Order for remainent injunction

1	2.	Venue for these matters is proper in this Court under 15 U.S.C. § 22 and 28 U.S.C.
2	§1391(b) and	d (c).
3	3.	The Complaint alleges that Endo engaged in violations of federal antitrust and
4	state antitrus	t and consumer protection laws by entering an agreement that foreclosed competition
5	from generic	equivalents of the brand-name drug Lidoderm [®] and later reduced competition
6	between selle	ers of generic lidocaine patches.
7	4.	Endo admits the facts necessary to establish the personal and subject matter
8	jurisdiction of	of this Court in this matter only.
9	5.	Endo denies the charges in the Complaint and disputes that the Party States are
10	entitled to ob	otain relief.
11	6.	This Order does not constitute any evidence against Endo, or an admission of
12	liability or w	rongdoing by Endo, in this case or in any other litigation. This Order shall not be
13	used in any v	way, as evidence or otherwise, in any other litigation or proceeding; provided,
14	<i>however</i> , tha	t this provision shall not prevent the Party States or Endo from using this Order in
15	any proceedi	ng regarding enforcement or modification of this ASO or as otherwise required by
16	law.	
17	7.	Entry of this ASO is in the public interest. The Party States and Endo have agreed
18	to stipulate to	o entry of this ASO to finally resolve the claims and litigation between them.
19		STIPULATIONS
20	1.	The Party States stipulate that they will not file litigation or any other proceedings
21	against Endo	asserting, or seeking remedies based on, Released Claims, other than any legal
22	proceeding r	egarding enforcement or modification of this ASO.
23	2.	Upon entry of this ASO by the Court, the Complaint shall be deemed dismissed
24	with prejudic	ce.
25		I. DEFINITIONS
26	IT IS	ORDERED that, as used in this ASO, the following definitions shall apply:
27	А.	"505(b)(2) Application" means an application filed with the United States Food
28	and Drug Ad	lministration pursuant to Section $505(b)(2)$ of the Federal Food, Drug and Cosmetic $\frac{3}{3}$
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1	Act, U.S.C. § 355(b)(2).
2	B. "ANDA" means an Abbreviated New Drug Application filed with the United States
3	Food and Drug Administration pursuant to Section 505(j) of the Federal Food, Drug and Cosmetic
4	Act, 21 U.S.C. § 355(j).
5	C. "Authorized Generic" means a Drug Product that is manufactured pursuant to an
6	NDA and Marketed in the United States under a name other than the proprietary name identified
7	in the NDA.
8	D. "Brand/Generic Settlement" means any agreement or understanding that settles a
9	Patent Infringement Claim in or affecting Commerce in the United States.
10	E. "Brand/Generic Settlement Agreement" means a written agreement that settles a
11	Patent Infringement Claim in or affecting Commerce in the United States.
12	F. "Branded Subject Drug Product" means a Subject Drug Product Marketed in the
13	United States under the proprietary name identified in the NDA for the Subject Drug Product.
14	G. "Commerce" has the same definition as it has in 15 U.S.C. § 44.
15	H. "Control" or "Controlled" means the holding of more than fifty percent (50%) of
16	the common voting stock or ordinary shares in, or the right to appoint more than fifty percent
17	(50%) of the directors of, or any other arrangement resulting in the right to direct the management
18	of, the said corporation, company, partnership, joint venture, or entity.
19	I. "Contingent Supply Agreement" means a Supply Agreement that: (i) is contingent
20	on the Generic Filer's inability to market the Generic Subject Drug Product on or after the
21	Generic Entry Date because (x) the FDA has not granted final approval of the Generic Filer's
22	ANDA or 505(b)(2) Application for the Generic Subject Drug Product and/or (y) the Generic
23	Filer cannot manufacture commercial quantities of the Generic Subject Drug Product; and (ii)
24	terminates within thirty (30) days after the Generic Filer has final FDA approval and can
25	manufacture commercial quantities of the Generic Subject Drug Product using good faith,
26	commercially reasonable efforts, provided, however, the Generic Filer may take delivery of,
27	Market, and sell quantities of Authorized Generic ordered prior to termination of the Supply
28	Agreement <i>so long as</i> the total quantity of Authorized Generic delivered to the Generic Filer 4
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following termination of the Supply Agreement: (i) does not exceed the total quantity needed by 1 the Generic Filer (as reflected in forecasts provided to the NDA Holder prior to termination of the 2 Supply Agreement) during the eight (8) months following (x) termination of the Supply 3 Agreement, if termination occurs after the Generic Entry Date, or (y) the Generic Entry Date, if 4 termination occurs before the Generic Entry Date; and (ii) is delivered within eight (8) months of 5 termination of the Supply Agreement. 6 J. "Drug Product" means a finished dosage form (e.g., tablet, capsule, solution, or 7 patch), as defined in 21 C.F.R. § 314.3(b), approved under a single NDA, ANDA or 505(b)(2) 8 9 Application, that contains a drug substance, generally, but not necessarily, in association with one or more other ingredients. 10 K. "Endo Pharmaceuticals" means Endo Pharmaceuticals Inc., any joint venture, 11 subsidiary, division, group, or affiliate Controlled currently or in the future by Endo 12 Pharmaceuticals Inc., their successors and assigns, and the respective directors, officers, 13 employees, agents, and representatives acting on behalf of each. 14 L. "Endo International" means Endo International plc, any joint venture, subsidiary, 15 16 division, group, or affiliate Controlled currently or in the future by Endo International plc, their successors and assigns, and the respective directors, officers, employees, agents, and 17 representatives acting on behalf of each. 18 M. "Endo" means Endo Pharmaceuticals and Endo International. 19 N. "Exception" means the following in a Brand/Generic Settlement: 20 compensation for saved future litigation expenses, *but only if* the total 1. 21 compensation the NDA Holder agrees to provide to the Generic Filer during the sixty (60) 22 day period starting thirty (30) days before and ending thirty (30) days after executing the 23 Brand/Generic Settlement Agreement does not exceed a maximum limit, which is initially 24 set at seven million, two hundred sixty-two thousand, eight hundred seventy-nine dollars 25 $(\$7,262,879)^1$ and shall be increased (or decreased) as of January 1 of each year following 26 27 For avoidance of doubt, this amount is intended to be equivalent to the maximum limit in the definition of Exception in the FTC Order, but is adjusted herein to reflect the 28 subsequent annual changes in the Producer Price Index as provided for in the FTC Order.

entry of this ASO by an amount equal to the percentage increase (or decrease) from the previous year in the annual average Producer Price Index for Legal Services (Series Id. PCU5411--5411--) published by the Bureau of Labor Statistics of the United States Department of Labor or its successor;

2. the right to Market, as of an agreed upon Generic Entry Date: (i) Generic Product(s) in the United States under an ANDA or 505(b)(2) Application (x) that is controlled by the Generic Filer and was not transferred to the Generic Filer by the NDA Holder, or (y) to which the Generic Filer has a license from a party other than the NDA Holder; or (ii) an Authorized Generic of the Subject Drug Product; this Exception 2 shall apply regardless of whether or not the Generic Filer must pay for the right to Market and, if so, the terms and conditions governing such payment;

3. provisions to facilitate, by means other than the transfer of goods or money, the Generic Filer's ability to secure or maintain final regulatory approval, or commence or continue the Marketing, of a Generic Product, by, *inter alia*, providing covenants, waivers, permissions, releases, dismissals of claims, and/or authorizations;

4. waiver or limitation of a claim for damages or other monetary relief based on prior Marketing of the Generic Subject Drug Product, *but only if* the NDA Holder and the Generic Filer do not agree, and have not agreed, to another Brand/Generic Settlement for a different Drug Product during the sixty (60) day period starting thirty (30) days before and ending thirty (30) days after the execution of the Brand/Generic Settlement Agreement; or

5. a continuation or renewal of a pre-existing agreement between an NDA Holder and a Generic Filer *but only if*: (i) the pre-existing agreement was entered into at least 90 days before the relevant Brand/Generic Settlement Agreement, (ii) the terms of the renewal or continuation, including the duration and the financial terms, are substantially similar to those in the pre-existing agreement, and (iii) entering into the continuation or renewal is not expressly contingent on agreeing to a Brand/Generic Settlement.

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1	O. "Exempted Agreement" means a Materials Agreement or Supply Agreement that
2	meets (x) all of the conditions in subparagraphs I.O.1-I.O.4 below, and (y) either (i) the conditions
3	in subparagraphs I.O.5 and I.O.6 below or (ii) the conditions in subparagraph I.O.7 below:
4	1. the price is above the Fully Allocated Manufacturing Cost, meaning:
5	a. if the Agreement is a Materials Agreement, the Materials Price charged
6	by the NDA Holder for Materials provided through the Materials Agreement is at
7	or above the Fully Allocated Manufacturing Cost incurred by the NDA Holder per
8	unit of the relevant Materials, or
9	b. if the Agreement is a Supply Agreement, the Supply Price charged by
10	the NDA Holder for the Authorized Generic of the Subject Drug Product is at or
11	above the Fully Allocated Manufacturing Cost incurred by the NDA Holder per
12	unit of the Authorized Generic of the Subject Drug Product provided under the
13	agreement;
14	2. the Brand/Generic Settlement Agreement containing or incorporating the
15	Materials Agreement or Supply Agreement is the only Brand/Generic Settlement
16	Agreement that the NDA Holder and the Generic Filer have entered, or agreed to enter,
17	during the sixty (60) day period starting thirty (30) days before and ending thirty (30)
18	days after the execution of the Brand/Generic Settlement Agreement;
19	3. within fourteen (14) days after signing the Brand/Generic Settlement Agreement
20	containing or incorporating the Materials Agreement or Supply Agreement, Endo
21	Submitted to the Monitor a full and complete copy of the Brand/Generic Settlement
22	Agreement, including any Materials Agreement and/or Supply Agreement;
23	4. within fourteen (14) days after the NDA Holder provides to the Generic Filer
24	the Materials Price or Supply Price, as applicable, Endo Submitted to the Monitor
25	notification of the relevant Materials Price or Supply Price;
26	5. within thirty (30) days after beginning supply under the relevant Materials
27	Agreement or Supply Agreement, the NDA Holder Submitted to the Monitor:
28	7
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1	a. if a Materials Agreement, a verified written statement containing (i) the
2	Fully Allocated Manufacturing Cost per unit for the Materials and (ii) a detailed
3	calculation of the Fully Allocated Manufacturing Cost for the Materials, stated
4	separately by cost component and on a per-unit basis; and
5	b. if a Supply Agreement, a verified written statement containing (i) the
6	Fully Allocated Manufacturing Cost per unit for the relevant Authorized Generic of
7	the Subject Drug Product and (ii) a detailed calculation of the Fully Allocated
8	Manufacturing Cost for the Authorized Generic of the Subject Drug Product, stated
9	separately by cost component and on a per-unit basis; and
10	6. if the NDA Holder is not Endo, the Materials Agreement or Supply Agreement,
11	as applicable, requires the NDA Holder to (i) provide the notification required by
12	subparagraphs I.O.5. and (ii) cooperate with any reasonable request by the Monitor or
13	staff of the FTC for documents and information to determine the relevant Fully Allocated
14	Manufacturing Cost, including without limitation and subject to any demonstrated legally
15	recognized privilege, providing the Monitor reasonable access to personnel, books,
16	documents, and records kept in the ordinary course of business;
17	7. notwithstanding subparagraph I.O.5. and subparagraph I.O.6., a Materials
18	Agreement or Supply Agreement in which Endo is the Generic Filer shall also be
19	considered an Exempted Agreement if it complies with subparagraphs I.O.1. to 4 and:
20	a. if a Materials Agreement, Endo Submits to the Monitor within thirty (30)
21	days of beginning to receive the Materials, a verified written statement containing
22	(i) Endo's best estimate of what would be the Fully Allocated Manufacturing Cost
23	per unit for the Materials if manufactured or sourced by the Generic Filer, including
24	a separate estimate of each cost component on a per-unit basis, and (ii) a
25	description of the terms and conditions of any agreement(s), offer(s), purchase
26	order(s), or price quote(s) Endo has entered into or received for supply of the
27	Materials in connection with manufacture of the Subject Drug Product and other
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1	facts and circumstances, if any, that Endo deems relevant to understanding such
2	terms and conditions; and
3	b. if a Supply Agreement, it is a Contingent Supply Agreement and Endo
4	Submits to the Monitor within thirty (30) days of beginning to receive the
5	Authorized Generic, a verified written statement containing (i) Endo's best estimate
6	of what would be the Fully Allocated Manufacturing Cost per unit for the Subject
7	Drug Product if manufactured by the Generic Filer and (ii) a detailed calculation of
8	the estimated Fully Allocated Manufacturing Cost, including an estimate of each
9	cost component on a per-unit basis.
10	P. "FTC" means the United States Federal Trade Commission.
11	Q. "FTC Order" means the Stipulated Order for Permanent Injunction entered in the
12	United States District Court for the Northern District of California, in Federal Trade Commission
13	v. Endo Pharmaceuticals Inc., Civ. Action No. 17-cv-00312 (Document 25, February 2, 2017).
14	R. "Fully Allocated Manufacturing Cost" means: (1) direct costs incurred to produce
15	or, if applicable, to acquire, the Subject Drug Product or Materials, determined in accordance with
16	GAAP, as consistently applied in accordance with past practice and in the ordinary course of
17	business, including, but not limited to (x) acquisition costs or (y) if applicable, materials, labor,
18	manufacturing costs, packaging, labeling, testing, quality control, storage, insurance, and product
19	maintenance; (2) the cost to ship the Subject Drug Product or Materials to the Generic Filer, and
20	(3) administrative and overhead expenses associated with production or, if applicable, the
21	acquisition of the Subject Drug Product or Materials, including, but not limited to, administrative
22	labor costs, maintenance, information technology, quality assurance, insurance, depreciation of the
23	equipment, and depreciation of the facility, allocated in accordance with past practice and in the
24	ordinary course of business. To the extent the NDA Holder does not allocate administrative and
25	overhead expenses associated with the Subject Drug Product to the Subject Drug Product, for
26	purposes of this ASO such administrative and overhead expenses shall be deemed to be a
27	proportion of the NDA Holder's COGS of the Subject Drug Product to the NDA Holder's total
28	COGS (for purposes of this definition, COGS means the NDA Holder's cost of goods sold, 9
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1	determined in accordance with GAAP, as consistently applied in accordance with past practice and
2	in the ordinary course of business).
3	S. "Generic Entry Date" means the date in a Brand/Generic Settlement Agreement,
4	whether certain or contingent, on or after which a Generic Filer is authorized by the NDA Holder
5	to begin manufacturing, using, importing or Marketing the Generic Subject Drug Product.
6	T. "Generic Filer" means a party to a Brand/Generic Settlement who controls an
7	ANDA or 505(b)(2) Application for the Subject Drug Product or has the exclusive right under
8	such ANDA or 505(b)(2) Application to distribute the Subject Drug Product.
9	U. "Generic Product" means a Drug Product manufactured and/or sold under an
10	ANDA or pursuant to 505(b)(2) Application.
11	V. "Generic Subject Drug Product" means the Generic Product that is the subject of
12	the Patent Infringement Claim being resolved by the Brand/Generic Settlement.
13	W. "Lidoderm Settlement Agreement" means the Settlement and License Agreement
14	between Endo Pharmaceuticals Inc. and Watson Laboratories, Inc. resolving the ANDA patent
15	litigation involving the brand-name drug Lidoderm [®] that is the subject of the Complaint in this
16	action.
17	X. "Market," "Marketed," or "Marketing" means the promotion, offering for sale,
18	sale, or distribution of a Drug Product.
19	Y. "Materials" means components or ingredients used in the manufacturing of a
20	Subject Drug Product, including, but not limited to, hard-to-source excipients, hard-to-source
21	active pharmaceutical ingredients, hard-to-source packaging, devices, or kits for injectables.
22	Z. "Materials Agreement" means provisions in, or incorporated into, a Brand/Generic
23	Settlement Agreement providing for the supply of Materials to the Generic Filer by the NDA
24	Holder for securing and/or maintaining regulatory approval, or manufacturing and Marketing by
25	the Generic Filer of the Subject Drug Product, including the terms and conditions of any such
26	supply.
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1	AA. "Materials Price" means the total actual per-unit price charged by the NDA Holder
2	for Materials provided through a Materials Agreement, including any transfer price and royalty to
3	be paid by the Generic Filer, net of any discounts, allowances, rebates, or other reductions.
4	BB. "Monitor" means an individual appointed pursuant to Paragraph IV of the FTC
5	Order.
6	CC. "NDA" means a New Drug Application filed with the United States Food and Drug
7	Administration pursuant to Section 505(b) of the Federal Food, Drug and Cosmetic Act, 21 U.S.C.
8	§ 355(b), including all changes or supplements thereto that do not result in the submission of a
9	new NDA.
10	DD. "NDA Holder" means a party to a Brand/Generic Settlement that controls the NDA
11	for the Subject Drug Product or has the exclusive right to distribute the Branded Subject Drug
12	Product in the United States.
13	EE. "No-AG Commitment" means any agreement with, or commitment or license to,
14	the Generic Filer that prohibits, prevents, restricts, requires a delay of, or imposes a condition
15	precedent upon the research, development, manufacture, regulatory approval, or Marketing of an
16	Authorized Generic, provided, however, that agreement by the Generic Filer to pay royalties to
17	the NDA Holder for the right to Market the Generic Subject Drug Product or an Authorized
18	Generic of the Subject Drug Product, including agreement on the terms and conditions governing
19	payment of such royalties, shall not be considered a No-AG Commitment.
20	FF. "Patent Infringement Claim" means any allegation threatened in writing or
21	included in a complaint filed with a court of law that a Generic Product may infringe one or more
22	U.S. Patents held by, or licensed to, an NDA Holder.
23	GG. "Payment by the NDA Holder to the Generic Filer" means a transfer of value,
24	other than a No-AG Commitment, by the NDA Holder to the Generic Filer (including, but not
25	limited to, money, goods, or services), regardless of whether the Generic Filer purportedly
26	transfers value in return, where such transfer is either (i) expressly contingent on entering a
27	Brand/Generic Settlement Agreement, or (ii) agreed to during the sixty (60) day period starting
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thirty (30) days before and ending thirty (30) days after executing a Brand/Generic Settlement
 Agreement.

3 HH. "Subject Drug Product" means the Drug Product for which one or more Patent
4 Infringement Claims are settled under a given Brand/Generic Settlement. For purposes of this
5 ASO, the Drug Product of the NDA Holder and the Generic Filer to the same Brand/Generic
6 Settlement shall be considered to be the same Subject Drug Product.

II. "Supply Agreement" means provisions in, or incorporated into, a Brand/Generic
Settlement Agreement providing for the supply of the Subject Drug Product to the Generic Filer
by the NDA Holder for the Marketing by the Generic Filer of an Authorized Generic on or after
the Generic Entry Date, including the terms and conditions of any such supply.

JJ. "Supply Price" means the total actual per-unit price charged by the NDA Holder
for supply provided through a Supply Agreement, including any transfer price and royalty to be
paid by the Generic Filer for the right to sell an Authorized Generic of the Subject Drug Product,
net of any discounts, allowances, rebates, or other reductions.

15 KK. "U.S. Patent" means any patent issued by the United States Patent and Trademark
16 Office, including all renewals, derivations, divisions, reissues, continuations, continuations-in
17 part, modifications, or extensions thereof.

18 LL. "Verified Written Report" means a report that must be submitted to the FTC
19 pursuant to Paragraph V of the FTC Order.

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II. RELEASED CLAIMS

IT IS FURTHER ORDERED that:

A. Release. The Group One States² and the Group Two States³ shall release and/or
stipulate and agree not to assert claims as follows:

(1) Group One States' Released Claims: In consideration of the injunctive

- provisions and Settlement Amount contained herein, Group One States will be
- 26
 27 Group One States include: Alabama, Iowa, Indiana, Maryland, Minnesota, Mississippi, Oklahoma, Utah, Virginia, Washington, and Wisconsin.
- 28 Group Two States include: Arkansas, Florida, Hawaii, Idaho, Illinois, Missouri,

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deemed, upon the Effective Date, to have fully, finally, and forever released Endo and affiliated entities and their current and former officers, directors, employees, agents, other associated persons and attorneys (collectively "Releasees") from any and all manner of claims, counterclaims, set-offs, demands, actions, rights, liabilities, costs, debts, expenses, attorneys' fees, and causes of action of any type, whether or not accrued in whole or in part, known or unknown, under federal or state law, arising from the Lidoderm Patent Settlement Agreement and from the facts, matters, transactions, events, occurrences, acts, disclosures, statements, omissions, or failures to act arising from said agreement ("Covered Conduct") that were asserted or alleged, or could have been asserted or alleged, in the Complaint by each Group One State or the Attorney General of each Group One State in any capacity, including but not limited his/her sovereign capacity as the chief law enforcement officer, parens patriae or proprietary capacity ("Group One States' Resolved Claims"). (2) Group Two States' Released Claims: In consideration of the injunctive provisions and Settlement Amount contained herein, Group Two States will be deemed, upon the Effective Date, to have fully, finally, and forever released Releasees from any and all manner of claims, counterclaims, set-offs, demands, actions, rights, liabilities, costs, debts, expenses, attorneys' fees, and causes of action of any type, whether or not accrued in whole or in part, known or unknown, under federal or state law, arising from the Covered Conduct that were asserted or alleged, or could have been asserted or alleged, in the Complaint by each Group Two State in a law enforcement capacity or the Attorney General of each Group Two State in his/her sovereign capacity as the chief law enforcement officer. In addition, each Group Two State is otherwise deemed to have stipulated and agreed that, in light of the terms of this ASO, neither it nor its Attorney General will assert any claim in any capacity or any claim on behalf of its departments, commissions, divisions, districts and other 13

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1	agencies, and the predecessors, successors, administrators and assigns of any
2	of the foregoing against Endo arising from the Covered Conduct that were
3	asserted or alleged, or could have been asserted or alleged, in its Complaint.
4	(All claims covered by this Section II.A.(2) "Group Two States' Resolved
5	Claims.)
6	The Group One States' Resolved Claims and the Group Two States' Resolved Claims
7	(together, the "Collective Resolved Claims") shall not include the Excluded Claims defined
8	below. In addition, in connection with the release provided above, the Party States expressly
9	waive, release, and forever discharge any and all provisions, rights, and benefits conferred by any
10	law of any state or territory of the United States or other jurisdiction, or principle of common law,
11	which is similar, comparable or equivalent to § 1542 of the California Civil Code, which reads:
12	Section 1542. General Release; extent. A general release does not extend to claims
13	which the creditor does not know or suspect to exist in his or her favor at the time of executing the release, which if known by him or her must have materially affected his or
14	her settlement with the debtor;
15	A Party State may hereafter discover facts other than or different from those which he, she, or it
16	knows or believes to be true with respect to the Collective Resolved Claims, but each Party State
17	hereby expressly waives and fully, finally, and forever settles, releases, and discharges, upon this
18	ASO becoming final, any known or unknown, suspected or unsuspected, asserted or unasserted,
19	contingent or non-contingent claim that would otherwise fall within the definition of Collective
20	Resolved Claims, whether or not concealed or hidden, without regard to the subsequent discovery
21	or existence of such different or additional facts, provided, however, that this shall not apply to
22	claims arising from documents or information intentionally withheld or concealed by Endo in
23	response to a subpoena served by a Party State in an effort to obstruct that Party State's
24	investigation. This provision shall not in any way expand the scope of the Collective Resolved
25	Claims and shall not convert what is a limited release into a general release.
26	B. Excluded Claims. The release provided herein shall not have an effect on the
27	following Excluded Claims and such Excluded Claims shall not be included within the Collective
28	Resolved Claims:
	Joint Motion for Entry of Stipulated Order for Permanent Injunction

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1	(1) claims or potential claims, under federal or state laws, of persons other than the
2	Party States ⁴ against Endo, including, but not limited to, any claims or
3	potential claims asserted in In re Lidoderm Antitrust Litigation, No. 14-md-
4	02521-WHO (N.D. Cal.);
5	(2) claims for alleged violations of state revenue codes;
6	(3) claims based on alleged breach of expressed or implied warranty claims or
7	other liability for defective or deficient products and services provided by
8	Endo;
9	(4) claims based on alleged failure to deliver goods or services due; or
10	(5) any claim for liability under common law, any statute, regulation, or rule
11	lawfully promulgated under each state's administrative code or other enabling
12	legislation for any conduct other than the Covered Conduct, including but not
13	limited to:
14	i. claims regarding Medicare or Medicaid fraud, false claims, unlawful
15	kickbacks, off-label marketing, unfair or deceptive representations,
16	false advertising or product liability;
17	ii. claims for criminal liability; or
18	iii. claims for breach of this ASO.
19	C. Res Judicata. The ASO shall be deemed to have rendered any of the Collective
20	Resolved Claims as res judicata.
21	III. SETTLEMENT PAYMENT
22	IT IS FURTHER ORDERED that:
23	A. Endo Pharmaceuticals Inc. shall make a payment to the Party States in the amount
24	of U.S. \$2,265,983.59 ("Settlement Amount") within the later of thirty (30) calendar days after
25	the Effective Date or thirty (30) calendar days after receiving written payment processing
26	instructions and any related documentation reasonably requested by Endo. The Settlement
27	⁴ For avoidance of doubt, this ASO does not release any claims that have been or
28	could be brought by private consumers, even if those claims could have been asserted by one or more of the Party States in their <i>parens patriae</i> capacity.
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Amount shall be apportioned among the Party States at the sole discretion of the Party States as may be agreed upon among them, and Endo shall have no responsibility for or influence with 2 respect to that allocation, which may include any or all of the following: (i) payment of 3 attorneys' fees and expenses; (ii) antitrust or consumer protection law enforcement; (iii) for 4 deposit into a state antitrust or consumer protection account (e.g., revolving account, trust 5 account), for use in accordance with the state laws governing that account; or (iv) for such other 6 purpose as the Attorneys General deem appropriate, consistent with the various states' laws.⁵ 7 The Party States shall provide Endo Pharmaceuticals Inc. with written payment processing 8 9 instructions for payment of the Settlement Amount by electronic transfer.

B. Endo Pharmaceuticals Inc. warrants that, as of the date of this Settlement 10 11 Agreement, it is not insolvent, nor will its Settlement Payment render it insolvent within the meaning of and/or for the purposes of the United States Bankruptcy Code. If (i) a case is 12 commenced with respect to Endo under Title 11 of the United States Code (Bankruptcy), or a 13 14 trustee, receiver or conservator is appointed under any similar law, and (ii) a court of competent jurisdiction enters a final order determining the Settlement Payment, or any portion thereof, by 15 16 or on behalf of Endo Pharmaceuticals Inc., to be a preference, voidable transfer, fraudulent transfer or similar transaction, and (iii) pursuant to an order of a court of competent jurisdiction 17 monies paid by Endo Pharmaceuticals Inc. pursuant to this Settlement Agreement are either not 18 delivered or are returned to Endo Pharmaceuticals Inc. or the trustee, receiver, or conservator 19 appointed by a court in any bankruptcy proceeding with respect to Endo Pharmaceuticals Inc., 20 21 then the releases given and judgment entered in favor of Endo pursuant to this Settlement 22 Agreement shall be null and void.

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IV. **PERMANENT INJUNCTION**

- **IT IS FURTHER ORDERED** that:
- A. In connection with any actions in or affecting Commerce, Endo shall cease and
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5 As deemed appropriate in the discretion of the Attorney General of Hawaii, 27 Hawaii's portion of the Settlement Amount may also be distributed to state agencies and entities whose programs provided funding, directly or indirectly, for Lidoderm and Lidocaine 28 prescriptions, which programs were not completely funded through general fund appropriations.

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1	desist from, either directly or indirectly, or through any corporate or other device, individually or
2	collectively entering into a Brand/Generic Settlement that includes:
3	1. (i) a No-AG Commitment and (ii) an agreement by the Generic Filer not to
4	research, develop, manufacture, or Market the Subject Drug Product for
5	any period of time; or
6	2. (i) any Payment by the NDA Holder to the Generic Filer that is not an
7	Exception or an Exempted Agreement and (ii) an agreement by the Generic
8	Filer not to research, develop, manufacture, or Market the Subject Drug
9	Product for any period of time,
10	provided, however, that any agreement entered into by an entity prior to that entity becoming part
11	of Endo is not subject to the terms of this ASO.
12	B. In the event of a material change in the law governing the antitrust implications of
13	Brand/Generic Settlements, the Party States will consider, in good faith, modifications to this
14	ASO proposed by Endo.
15	C. Nothing in this ASO shall prohibit Endo from entering a written agreement,
16	including a Brand/Generic Settlement, for which Endo has submitted to the State Reporting and
17	Enforcement Council ("SREC"), by written communication through the Liaison States (defined
18	below), a request for prior approval of the agreement, which shall include any information
19	provided to the FTC pursuant to Paragraph III of the FTC Order, so long as (1) within thirty (30)
20	days of the Liaison States' receipt of the request for prior approval under this provision, the
21	Liaison States have not notified Endo in writing that, after considering the request in good faith,
22	the SREC believes the relevant agreement raises substantial questions regarding violation of state
23	or federal antitrust law or any other applicable law that the Party States have authority to enforce
24	and of the reasons for such a belief, or (2) Endo has received, in a written communication by the
25	Liaison States, the prior approval of the SREC.
26	D. Nothing in this Section IV shall prohibit Endo from executing a written agreement
27	so long as such agreement contains a provision or provisions expressly stating: (1) Endo will
28	submit to the SREC by written communication through the Liaison States, a request for prior 17

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approval of the agreement, and (2) the agreement is not effective, and shall not become effective,
until and unless (i) thirty (30) days have passed since the request for prior approval was submitted
to the Liaison States and the Liaison States have not notified Endo in writing that the SREC
believes the agreement raises substantial questions regarding violation of state or federal antitrust
law or any other applicable law that the Party States have authority to enforce, or (ii) the SREC
has approved of the agreement, as reflected in a written communication to Endo from the Liaison
States.

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V. STATES REPORTING AND ENFORCEMENT COUNCIL IT IS FURTHER ORDERED that:

A. The Party States shall have authority for enforcing this ASO as set forth herein. 10 11 B. The Party States shall establish a States Reporting and Enforcement Council ("SREC") consisting initially of the States of Mississippi and Minnesota. Any Party State may 12 resign from or join the SREC upon 10 days' prior written notice to Endo. The Party States shall 13 exercise their rights under this ASO solely through the SREC, which shall have sole and 14 exclusive right and authority, among other things, to seek to enforce this ASO, to implement the 15 16 reporting requirements and receive from Endo the materials specified in Section VI herein (Reporting and Monitoring Obligations), and to exercise any rights to access information under 17 Section VIII (Access to Information). 18

C. The SREC shall designate up to five (5) of its members to serve as the Liaison 19 States. The initial Liaison States shall be Mississippi and Minnesota. The members serving as 20 21 Liaison States may be changed from time to time at the discretion of the SREC, but in any event, the SREC shall notify Endo within three (3) business days of the selection of any initial and 22 subsequent Liaison States, and shall provide the names and contact information for the authorized 23 representatives of any Liaison States. The SREC shall communicate with Endo, and Endo shall 24 communicate with the SREC, solely through the Liaison States concerning any matter covered by 25 26 or relating to this ASO, and Endo may rely exclusively on such communications with the Liaison States as being on behalf of and having the full authority of the SREC. By way of clarification 27 and not limitation, Endo shall be obligated to provide only to the Liaison States any reports, 28

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information, or notices required under this ASO, which Endo shall transmit by electronic mail
and/or overnight delivery service and which shall be effective upon transmission. The members
of the SREC and the Liaison States, as of the Effective Date, are identified on Exhibit A hereto
along with contact information for their respective representatives. All notices or
communications to Endo under this ASO shall be made to the individuals identified on Exhibit B
hereto, as may be amended in writing from time to time, and shall be made by electronic mail
and/or by overnight delivery service.

B. The SREC shall have the sole and exclusive authority on behalf of the Party States
to enter into and implement the terms of this ASO, as described herein. The SREC shall also
have the sole and exclusive authority on behalf of the Party States to seek to enforce this ASO, as
provided herein.

E. If the SREC believes that Endo is not in compliance with the terms of this ASO, 12 the SREC, through the Liaison States, shall give Endo written notice of such alleged non-13 compliance and the reasons why the SREC believes that Endo is not complying with this ASO. 14 Endo shall have twenty (20) business days from the date of receipt of such notice to respond in 15 16 writing unless otherwise agreed by the Parties. If the SREC is not satisfied with Endo's response, the Liaison States shall so notify Endo in writing, and Endo shall have seventy-five (75) calendar 17 days from the date of receipt of such notice to cure such alleged non-compliance. If after such 18 time, the SREC believes that Endo remains not in compliance with this ASO, the SREC may seek 19 to enforce this ASO with the Court in a civil enforcement proceeding, seeking such relief as is 20 available under applicable laws, including, if so available, fees and costs thereof, and civil 21 penalties. Such civil enforcement proceedings shall be governed by a preponderance of the 22 evidence standard, absent an applicable statutory provision or other binding legal authority to the 23 contrary. 24

F. In the event that the SREC believes that Endo remains in non-compliance with any of the provisions of this ASO other than those in Section IV (Permanent Injunction) following the notice and cure period provided in Section V.E., above, the SREC may seek to enforce such provisions of this ASO by filing a motion in this Court, seeking such relief as is available under

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applicable laws, including, if so available, fees and costs thereof, and civil penalties. 1 G. In the event that the SREC believes that Endo remains in non-compliance with any 2 of the provisions of Section IV (Permanent Injunction) of this ASO following the notice and cure 3 period provided in Section V.E., above, the SREC shall request that the FTC review Endo's 4 compliance with the corresponding provisions of Paragraph II of the FTC Order and consider 5 pursuing coordinated enforcement proceedings under this ASO and the FTC Order. In the event 6 that the FTC declines or fails to agree after forty-five (45) days from such a request by the SREC 7 to take action to enforce Paragraph II of the FTC Order, the SREC may then seek to enforce 8 9 Section IV of this ASO by filing a motion in this Court. The SREC, through the Liaison States, shall provide written notice to Endo within three (3) business days of any request(s) to the FTC 10 provided herein. 11 H. Nothing in this Section V, however, prohibits the SREC from pursuing immediate 12 enforcement for alleged non-compliance with this ASO upon a showing that delay would cause 13 irreparable harm or would prevent the SREC from seeking adequate enforcement of the ASO. 14 **REPORTING AND MONITORING OBLIGATIONS** VI. 15 IT IS FURTHER ORDERED that: 16 A. Each Verified Written Report that Endo is required to submit to the FTC shall also 17 be submitted to the SREC by service upon the Liaison States within five (5) business days of 18 submission to the FTC. 19 B. Each Branded/Generic Settlement Agreement submitted to the FTC pursuant to 20 21 Section 1112(a) of the Medicare Prescription Drug, Improvement, and Modernization Act of 22 2003 shall also be submitted to the SREC by service upon the Liaison States within five (5) business days of submission to the FTC. 23 C. Endo shall consent to the FTC providing the SREC with a copy of each written 24 report that the Monitor is required to submit to the FTC under the FTC Order. 25 D. Endo shall consent to the FTC providing the SREC with copies of any and all of 26 its submissions to the FTC under the FTC Order made between February 2, 2017 and the entry of 27 this ASO ("FTC Submissions"), provided that: (a) Endo shall have five (5) business days after 28

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1 the entry of this ASO to notify the necessary third parties of the pending disclosure; (b) such third parties shall have twenty (20) business days from the entry of this ASO to assert an objection to 2 this Court; and (c) Endo's consent shall not be effective until, either: (i) twenty-six (26) business 3 days from the entry of this ASO have passed without a third party having asserted an objection to 4 this Court; or (ii) if a third party has asserted an objection, this Court has issued a ruling 5 6 permitting such disclosure over any such objection(s). If, after this consent is effective, the SREC has requested the FTC Submissions from the FTC, but the FTC has not been willing to provide 7 them to the SREC, Endo will produce the FTC Submissions to the SREC. 8

E. Except as provided in Section VI.F. below, any agreements, information, or 9 documents submitted by Endo pursuant to this ASO ("Endo Materials") may be divulged or 10 11 disclosed by any Party State only to a person or entity who is an authorized representative or retained consultant or expert of any Party State, who is engaged in matters pertaining to the 12 implementation or enforcement of this ASO, and who has agreed to be bound by the provisions of 13 this Section VI.D. To the extent any Party State retains private counsel to investigate or assert 14 potential claims against Endo in matters that are not related to the implementation or enforcement 15 16 of this ASO, then no Endo Materials may be disclosed or divulged to those private counsel.

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F.

Notwithstanding the foregoing Section VI.D,

Endo Materials may be disclosed in the course of a legal proceeding
 between Parties to this ASO to enforce or modify this ASO, provided all parties to
 such a proceeding take reasonable steps to prevent disclosure of such Endo
 Materials to persons or entities who are not Parties to this ASO and Endo is given
 at least ten (10) business days' advance written notice and a reasonable
 opportunity to preserve the confidentiality of Endo Materials sought to be
 disclosed;

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 2. A Party State may disclose Endo Materials in response to a discovery
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 26 request or other legal process in a legal proceeding to which the Party State is a
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 28 party, or as otherwise required by law (other than a grand jury proceeding),
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such Endo Materials by proceeding under any applicable protective order and/or utilizing sealing procedures provided by law or court rule and by providing Endo with at least ten (10) business days' advance written notice and a reasonable opportunity to preserve the confidentiality of Endo Materials sought to be disclosed before disclosing such Endo Materials to a third party; and

3. Endo Materials may be disclosed to the FTC or the United States Department of Justice, provided Endo is given at least ten (10) business days' advance written notice of such disclosure.

The Party States acknowledge and agree that (i) the production of Endo Materials to 9 F. the Party States is compelled by the terms of this ASO; (ii) Endo may mark any such Endo 10 11 Materials, where appropriate, as trade secrets or otherwise exempt from public disclosure ("Confidential Endo Materials"); (iii) Endo does not waive any claimed exemption from public 12 disclosure under any law with respect to Confidential Endo Materials; and (iv) nothing in this 13 ASO shall prevent Endo from seeking appropriate relief under the law of any Party State 14 concerning public disclosures. In the event of a request for production of any Endo Materials 15 16 pursuant to any law regarding public disclosure of documents in the possession of a Party State (a "Records Request Notice"), such Party State, by and through its respective Attorney General, to 17 the extent not prohibited by law, shall (a) promptly provide Endo with notice and a copy of the 18 Records Request Notice, (b) designate any Confidential Endo Materials as falling within any 19 trade secret or similar exemptions from disclosure under that Party State's applicable laws, and 20 21 (c) notify Endo in advance of any disclosure of any Endo Materials in order to afford Endo the ability to seek appropriate relief under the Party State's law with respect to the disclosure of such 22 Endo Materials. 23

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G. An enforcement action in this Court pursuant to this ASO shall be the exclusive remedy for violation of this ASO. Neither the terms of this ASO nor any reports or notices 25 provided by Endo under this ASO shall operate as a waiver of any future claims by any third 26 party. Further, any such reports or notices provided by Endo to members of the SREC under this 27 ASO shall not be deemed to constitute actual or constructive notice of any claims as to any third 28 22

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1	party that has not received such reports or notices. No provision in this ASO may be used as
2	evidence by Endo or by the Party States in a proceeding other than an enforcement action
3	pursuant to this ASO.
4	VII. PROPOSED CHANGE OF CORPORATE CONTROL
5	IT IS FURTHER ORDERED that:
6	A. Endo shall notify the SREC at least thirty (30) days prior to:
7	1. Any proposed dissolution of Endo; or
8	2. Any proposed acquisition, merger, or consolidation of Endo; or
9	3. Any other change in Endo, including, but not limited to, assignment and the
10	creation, sale or dissolution of subsidiaries, if such change might affect the
11	compliance obligations arising out of this ASO.
12	B. Endo shall notify the SREC within ten (10) days after any filing by Endo of a petition
13	in bankruptcy.
14	C. Endo shall submit any notice required under this paragraph to the SREC by service
15	upon the Liaison States.
16	VIII. ACCESS TO INFORMATION
17	IT IS FURTHER ORDERED that:
18	A. For the purpose of determining or securing compliance with this ASO, subject to and
19	without limiting any legally recognized privilege, and upon written request with reasonable
20	advance notice, Endo shall:
21	1. timely respond to and cooperate with the SREC's reasonable request for
22	production of documents or information related to compliance, including
23	permitting any duly authorized representative of the SREC to access, during
24	office hours and in the presence of counsel, all facilities and access to inspect
25	and copy all non-privileged business records and documentary material (which
25 26	and copy all non-privileged business records and documentary material (which may be redacted for privilege) related to compliance with this ASO, including
26	may be redacted for privilege) related to compliance with this ASO, including

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1	memoranda, written justifications, economic models, and other records and
2	documents (in whatever form such records and documents are kept) in the
3	possession or under the control of Endo, which copying services shall be
4	provided by Endo in a timely manner at the request of the authorized
5	representative(s) of the SREC;
6	2. permit any duly authorized representative of the SREC to interview officers,
7	directors, or employees of Endo, who may have counsel present, regarding any
8	such matters; and
9	3. agree to accept service of process of any motion filed with the Court hereunder
10	and, subject to any objections Endo may assert, cooperate with any subpoenas
11	issued by a Party State in connection with any motion to enforce this ASO.
12	B. The SREC shall to the fullest extent possible coordinate any requests for information
13	under this Section with the FTC and Department of Justice, if involved. The SREC shall make
14	any and all requests for information hereunder solely through the Liaison States.
15	IX. EFFECT OF SETTLEMENTS WITH OTHER STATES
16	IT IS FURTHER ORDERED that:
	A. The Party States shall permit California to join the enforcement and reporting
17	in the rank same share content of Jen the encounter and reporting
17 18	provisions of this multistate ASO upon the submission of an amended California ASO by
18	provisions of this multistate ASO upon the submission of an amended California ASO by
18 19	provisions of this multistate ASO upon the submission of an amended California ASO by California and Endo reflecting that California's enforcement authority and each of Endo's
18 19 20	provisions of this multistate ASO upon the submission of an amended California ASO by California and Endo reflecting that California's enforcement authority and each of Endo's notification and reporting obligations, including but not limited to the provisions reflected in
18 19 20 21	provisions of this multistate ASO upon the submission of an amended California ASO by California and Endo reflecting that California's enforcement authority and each of Endo's notification and reporting obligations, including but not limited to the provisions reflected in Paragraphs IV.B, IV.C, and V through VIII (with the exception of VI.F regarding confidentiality)
18 19 20 21 22	provisions of this multistate ASO upon the submission of an amended California ASO by California and Endo reflecting that California's enforcement authority and each of Endo's notification and reporting obligations, including but not limited to the provisions reflected in Paragraphs IV.B, IV.C, and V through VIII (with the exception of VI.F regarding confidentiality) of the California ASO, shall be replaced by the corresponding provisions of this ASO.
 18 19 20 21 22 23 	provisions of this multistate ASO upon the submission of an amended California ASO by California and Endo reflecting that California's enforcement authority and each of Endo's notification and reporting obligations, including but not limited to the provisions reflected in Paragraphs IV.B, IV.C, and V through VIII (with the exception of VI.F regarding confidentiality) of the California ASO, shall be replaced by the corresponding provisions of this ASO. B. If following the filing or entry of this ASO, Endo enters into a similar settlement
 18 19 20 21 22 23 24 	provisions of this multistate ASO upon the submission of an amended California ASO by California and Endo reflecting that California's enforcement authority and each of Endo's notification and reporting obligations, including but not limited to the provisions reflected in Paragraphs IV.B, IV.C, and V through VIII (with the exception of VI.F regarding confidentiality) of the California ASO, shall be replaced by the corresponding provisions of this ASO. B. If following the filing or entry of this ASO, Endo enters into a similar settlement agreement with another state, or a series of similar agreements with individual states, the Party
 18 19 20 21 22 23 24 25 	provisions of this multistate ASO upon the submission of an amended California ASO by California and Endo reflecting that California's enforcement authority and each of Endo's notification and reporting obligations, including but not limited to the provisions reflected in Paragraphs IV.B, IV.C, and V through VIII (with the exception of VI.F regarding confidentiality) of the California ASO, shall be replaced by the corresponding provisions of this ASO. B. If following the filing or entry of this ASO, Endo enters into a similar settlement agreement with another state, or a series of similar agreements with individual states, the Party States shall coordinate their enforcement actions, and Endo's reporting obligations with such
 18 19 20 21 22 23 24 25 26 	provisions of this multistate ASO upon the submission of an amended California ASO by California and Endo reflecting that California's enforcement authority and each of Endo's notification and reporting obligations, including but not limited to the provisions reflected in Paragraphs IV.B, IV.C, and V through VIII (with the exception of VI.F regarding confidentiality) of the California ASO, shall be replaced by the corresponding provisions of this ASO. B. If following the filing or entry of this ASO, Endo enters into a similar settlement agreement with another state, or a series of similar agreements with individual states, the Party States shall coordinate their enforcement actions, and Endo's reporting obligations with such other state(s). The Parties will use their best efforts to negotiate an agreement among and

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1	settlement agreement(s) and Endo and the Party States will submit any such agreement to this
2	Court as an amendment to this ASO.
3	X. JURISDICTION
4	IT IS FURTHER ORDERED that this Court shall retain jurisdiction over these matters
5	for purposes of construction, modification, and enforcement of this ASO.
6	XI. TERMINATION
7	IT IS FURTHER ORDERED that this ASO shall terminate on February 2, 2027.
8	XII. DISMISSAL WITH PREJUDICE
9	IT IS FURTHER ORDERED that the Complaint shall be and hereby is dismissed with
10	prejudice. Each party to bear its own costs aside from the Settlement Amount paid to the Party
11	States.
12	SO ORDERED this <u>6th</u> day of <u>August</u> , 2019
13	Mafine M. Cherney
14	MAXINE M. CHESNEY
15	UNITED STATES DISTRICT JUDGE
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28	25
	Joint Motion for Entry of Stipulated Order for Permanent Injunction

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FOR ENDO PHARMACEUTICALS INC.:

By: 11 Arell: Name: 5 Title:

7/15/19 Date:

FOR ENDO INTERNATIONAL PLC:

By nyonell. Name: 050 Title:

By:

Date: ()

Date: 7/15/2019

George G. Gordon Dechert LLP COUNSEL FOR ENDO PHARMACEUTICALS INC. AND ENDO INTERNATIONAL PLC

EXHIBIT A Contact Information for Liaison States and SREC

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EXHIBIT B Contact Information for Endo Pharmaceuticals Inc. and Endo International plc

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