STATE OF NEW YORK

4370

2021-2022 Regular Sessions

IN SENATE

February 3, 2021

Introduced by Sen. BIAGGI -- read twice and ordered printed, and when printed to be committed to the Committee on Health

AN ACT to amend the public health law, in relation to preserving access to affordable drugs

The People of the State of New York, represented in Senate and Assembly, do enact as follows:

1	Section 1. Article 2-A of the public health law is amended by adding a
2	new title IV to read as follows:
3	TITLE IV
4	PRESERVING ACCESS TO AFFORDABLE DRUGS
5	Section 282. Definitions.
6	283. Preserving access to affordable drugs.
7	<u>§ 282. Definitions. For the purposes of this title, the following</u>
8	terms shall have the following meanings:
9	1. "ANDA" shall mean abbreviated new drug application.
10	2. "ANDA filer" shall mean a party that owns or controls an ANDA filed
11	with the federal food and drug administration or has the exclusive
12	rights under that ANDA to distribute the ANDA product.
13	3. "Agreement" shall mean anything that would constitute an agreement
14	<u>under state law.</u>
15	4. "Agreement resolving or settling a patent infringement claim"
16	includes any agreement that is entered into within thirty days of the
17	resolution or the settlement of the claim, or any other agreement that
18	is contingent upon, provides a contingent condition for, or is otherwise
19	related to the resolution or settlement of the claim. This shall
20	include, but is not limited to, the following:
21	(a) Any agreement required to be provided to the federal trade commis-
22	sion or the antitrust division of the United States Department of
23	Justice under the Medicare Prescription Drug, Improvement, and Modern-
24	<u>ization Act of 2003, Pub. L. No. 108-173;</u>

EXPLANATION--Matter in <u>italics</u> (underscored) is new; matter in brackets [-] is old law to be omitted.

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S. 4370

1	(b) Any agreement between a biosimilar or interchangeable product
2	applicant and a reference product sponsor under the Biologics Price
3	Competition and Innovation Act of 2009, Pub. L. No. 111-148, that
4	resolves patent claims between the applicant and sponsor.
5	5. "Biosimilar biological product application filer" shall mean a
б	party that owns or controls a biosimilar biological product application
7	filed with the federal food and drug administration pursuant to section
8	351(k) of the Public Health Service Act, 42 U.S.C. 262(k), for licensure
9	of a biological product as biosimilar to, or interchangeable with, a
10	reference product, or that has the exclusive rights under the applica-
11	tion to distribute the biosimilar biological product.
12	6. "NDA" shall mean a new drug application.
13	7. "Nonreference drug filer" shall mean either:
14	(a) An ANDA filer; or
15	(b) A biosimilar biological product application filer.
16	8. "Nonreference drug product" shall mean the product to be manufac-
17	tured under an ANDA that is the subject of the patent infringement
18	claim, a biosimilar biological product that is the product to be manu-
19	factured under the biosimilar biological product application that is the
20	subject of the patent infringement claim, or both.
21	9. "Patent infringement" shall mean infringement of any patent or of
22	any filed patent application, extension, reissue, renewal, division,
23	continuation, continuation in part, reexamination, patent term restora-
24	tion, patents of addition, and extensions thereof.
25	10. "Patent infringement claim" shall mean any allegation made to a
26	nonreference drug filer, whether or not included in a complaint filed
27	with a court of law, that its nonreference drug product or application
28	infringes any patent held by, or exclusively licensed to, the reference
29	<u>drug holder.</u>
30	11 "Defenence drug belder" shall mean either.
30	<u>11. "Reference drug holder" shall mean either:</u>
30 31	(a) A brand holder that is any of the following:
31	(a) A brand holder that is any of the following:
31 32	(a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed
31 32 33	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21
31 32 33 34	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b); (ii) A person owning or controlling enforcement of the patent listed in the approved drug products with therapeutic equivalence evaluations
31 32 33 34 35	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b); (ii) A person owning or controlling enforcement of the patent listed
31 32 33 34 35 36	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b); (ii) A person owning or controlling enforcement of the patent listed in the approved drug products with therapeutic equivalence evaluations in connection with the NDA; or (iii) The predecessors, subsidiaries, divisions, groups, and affil-
31 32 33 34 35 36 37	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b); (ii) A person owning or controlling enforcement of the patent listed in the approved drug products with therapeutic equivalence evaluations in connection with the NDA; or (iii) The predecessors, subsidiaries, divisions, groups, and affiliates controlled by, controlling, or under common control with, any of
31 32 33 34 35 36 37 38	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b); (ii) A person owning or controlling enforcement of the patent listed in the approved drug products with therapeutic equivalence evaluations in connection with the NDA; or (iii) The predecessors, subsidiaries, divisions, groups, and affiliates controlled by, controlling, or under common control with, any of the entities described in subparagraph (i) or (ii) of this paragraph,
31 32 33 34 35 36 37 38 39 40 41	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b); (ii) A person owning or controlling enforcement of the patent listed in the approved drug products with therapeutic equivalence evaluations in connection with the NDA; or (iii) The predecessors, subsidiaries, divisions, groups, and affiliates controlled by, controlling, or under common control with, any of the entities described in subparagraph (i) or (ii) of this paragraph, with control to be presumed by direct or indirect share ownership of
31 32 33 34 35 36 37 38 39 40 41 42	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b); (ii) A person owning or controlling enforcement of the patent listed in the approved drug products with therapeutic equivalence evaluations in connection with the NDA; or (iii) The predecessors, subsidiaries, divisions, groups, and affiliates controlled by, controlling, or under common control with, any of the entities described in subparagraph (i) or (ii) of this paragraph, with control to be presumed by direct or indirect share ownership of fifty percent or greater, as well as the licensees, licensors, succes-
31 32 33 34 35 36 37 38 39 40 41 42 43	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b); (ii) A person owning or controlling enforcement of the patent listed in the approved drug products with therapeutic equivalence evaluations in connection with the NDA; or (iii) The predecessors, subsidiaries, divisions, groups, and affiliates controlled by, controlling, or under common control with, any of the entities described in subparagraph (i) or (ii) of this paragraph, with control to be presumed by direct or indirect share ownership of fifty percent or greater, as well as the licensees, licensors, successors, and assigns of each of those entities; or
31 32 33 34 35 36 37 38 39 40 41 42 43 44	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b); (ii) A person owning or controlling enforcement of the patent listed in the approved drug products with therapeutic equivalence evaluations in connection with the NDA; or (iii) The predecessors, subsidiaries, divisions, groups, and affiliates controlled by, controlling, or under common control with, any of the entities described in subparagraph (i) or (ii) of this paragraph, with control to be presumed by direct or indirect share ownership of fifty percent or greater, as well as the licensees, licensors, successors, and assigns of each of those entities; or (b) A biological product license holder, which shall mean any of the
31 32 33 34 35 36 37 38 39 40 41 42 43 44 45	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b); (ii) A person owning or controlling enforcement of the patent listed in the approved drug products with therapeutic equivalence evaluations in connection with the NDA; or (iii) The predecessors, subsidiaries, divisions, groups, and affiliates controlled by, controlling, or under common control with, any of the entities described in subparagraph (i) or (ii) of this paragraph, with control to be presumed by direct or indirect share ownership of fifty percent or greater, as well as the licensees, licensors, successors, and assigns of each of those entities; or (b) A biological product license holder, which shall mean any of the following:
31 32 33 34 35 36 37 38 39 40 41 42 43 44 45 46	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b); (ii) A person owning or controlling enforcement of the patent listed in the approved drug products with therapeutic equivalence evaluations in connection with the NDA; or (iii) The predecessors, subsidiaries, divisions, groups, and affiliates controlled by, controlling, or under common control with, any of the entities described in subparagraph (i) or (ii) of this paragraph, with control to be presumed by direct or indirect share ownership of fifty percent or greater, as well as the licensees, licensors, successors, and assigns of each of those entities; or (b) A biological product license holder, which shall mean any of the following: (i) The holder of an approved biological product license application
31 32 33 34 35 36 37 38 39 40 41 423 442 45 46 47	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b); (ii) A person owning or controlling enforcement of the patent listed in the approved drug products with therapeutic equivalence evaluations in connection with the NDA; or (iii) The predecessors, subsidiaries, divisions, groups, and affiliates controlled by, controlling, or under common control with, any of the entities described in subparagraph (i) or (ii) of this paragraph, with control to be presumed by direct or indirect share ownership of fifty percent or greater, as well as the licensees, licensors, successors, and assigns of each of those entities; or (b) A biological product license holder, which shall mean any of the following: (i) The holder of an approved biological product license application for a biological drug product under section 351(a) of the Public Health
31 32 33 34 35 36 37 38 39 40 41 42 43 44 45 46 47 48	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b); (ii) A person owning or controlling enforcement of the patent listed in the approved drug products with therapeutic equivalence evaluations in connection with the NDA; or (iii) The predecessors, subsidiaries, divisions, groups, and affiliates controlled by, controlling, or under common control with, any of the entities described in subparagraph (i) or (ii) of this paragraph, with control to be presumed by direct or indirect share ownership of fifty percent or greater, as well as the licensees, licensors, successors, and assigns of each of those entities; or (b) A biological product license holder, which shall mean any of the following: (i) The holder of an approved biological product license application for a biological drug product under section 351(a) of the Public Health Service Act, 42 U.S.C. 262(a);
31 32 33 34 35 36 37 38 40 41 42 43 445 465 47 48 49	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b); (ii) A person owning or controlling enforcement of the patent listed in the approved drug products with therapeutic equivalence evaluations in connection with the NDA; or (iii) The predecessors, subsidiaries, divisions, groups, and affiliates controlled by, controlling, or under common control with, any of the entities described in subparagraph (i) or (ii) of this paragraph, with control to be presumed by direct or indirect share ownership of fifty percent or greater, as well as the licensees, licensors, successors, and assigns of each of those entities; or (b) A biological product license holder, which shall mean any of the following: (i) The holder of an approved biological product license application for a biological drug product under section 351(a) of the Public Health Service Act, 42 U.S.C. 262(a); (ii) A person owning or controlling enforcement of any patents that
31 32 33 34 35 36 37 38 40 41 42 43 45 46 47 48 49 50	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b): (ii) A person owning or controlling enforcement of the patent listed in the approved drug products with therapeutic equivalence evaluations in connection with the NDA; or (iii) The predecessors, subsidiaries, divisions, groups, and affiliates controlled by, controlling, or under common control with, any of the entities described in subparagraph (i) or (ii) of this paragraph, with control to be presumed by direct or indirect share ownership of fifty percent or greater, as well as the licensees, licensors, successors, and assigns of each of those entities; or (b) A biological product license holder, which shall mean any of the following: (i) The holder of an approved biological product license application for a biological drug product under section 351(a) of the Public Health Service Act, 42 U.S.C. 262(a); (ii) A person owning or controlling enforcement of any patents that claim the biological product that is the subject of the approved biological
31 32 33 34 35 36 37 39 41 42 43 45 46 47 489 50 51	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b); (ii) A person owning or controlling enforcement of the patent listed in the approved drug products with therapeutic equivalence evaluations in connection with the NDA; or (iii) The predecessors, subsidiaries, divisions, groups, and affiliates controlled by, controlling, or under common control with, any of the entities described in subparagraph (i) or (ii) of this paragraph, with control to be presumed by direct or indirect share ownership of fifty percent or greater, as well as the licensees, licensors, successors, and assigns of each of those entities; or (b) A biological product license holder, which shall mean any of the following: (i) The holder of an approved biological product license application for a biological drug product under section 351(a) of the Public Health Service Act, 42 U.S.C. 262(a); (ii) A person owning or controlling enforcement of any patents that claim the biological product that is the subject of the approved biological patent license application; or
31 32 33 34 35 36 37 39 401 423 445 46 47 489 501 512	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b); (ii) A person owning or controlling enforcement of the patent listed in the approved drug products with therapeutic equivalence evaluations in connection with the NDA; or (iii) The predecessors, subsidiaries, divisions, groups, and affiliates controlled by, controlling, or under common control with, any of the entities described in subparagraph (i) or (ii) of this paragraph, with control to be presumed by direct or indirect share ownership of fifty percent or greater, as well as the licensees, licensors, successors, and assigns of each of those entities; or (b) A biological product license holder, which shall mean any of the following: (i) The holder of an approved biological product license application for a biological drug product under section 351(a) of the Public Health Service Act, 42 U.S.C. 262(a); (ii) A person owning or controlling enforcement of any patents that claim the biological product that is the subject of the approved biological patent license application; or
31 32 33 35 36 37 39 412 43 45 46 47 490 512 53	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b): (ii) A person owning or controlling enforcement of the patent listed in the approved drug products with therapeutic equivalence evaluations in connection with the NDA; or (iii) The predecessors, subsidiaries, divisions, groups, and affiliates controlled by, controlling, or under common control with, any of the entities described in subparagraph (i) or (ii) of this paragraph, with control to be presumed by direct or indirect share ownership of fifty percent or greater, as well as the licensees, licensors, successors, and assigns of each of those entities; or (b) A biological product license holder, which shall mean any of the following: (i) The holder of an approved biological product license application for a biological drug product under section 351(a) of the Public Health Service Act, 42 U.S.C. 262(a); (ii) A person owning or controlling enforcement of any patents that claim the biological product that is the subject of the approved biological patent license application; or (iii) The predecessors, subsidiaries, divisions, groups, and affiliates controlled by, controlling, or under common control with, any of
31 32 33 34 35 36 37 39 401 423 445 46 47 489 501 512	 (a) A brand holder that is any of the following: (i) The holder of an approved NDA for a drug product application filed under section 505(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(b); (ii) A person owning or controlling enforcement of the patent listed in the approved drug products with therapeutic equivalence evaluations in connection with the NDA; or (iii) The predecessors, subsidiaries, divisions, groups, and affiliates controlled by, controlling, or under common control with, any of the entities described in subparagraph (i) or (ii) of this paragraph, with control to be presumed by direct or indirect share ownership of fifty percent or greater, as well as the licensees, licensors, successors, and assigns of each of those entities; or (b) A biological product license holder, which shall mean any of the following: (i) The holder of an approved biological product license application for a biological drug product under section 351(a) of the Public Health Service Act, 42 U.S.C. 262(a); (ii) A person owning or controlling enforcement of any patents that claim the biological product that is the subject of the approved biological patent license application; or

S. 4370

1	fifty percent or greater, as well as the licensees, licensors, succes-
2	sors, and assigns of each of those entities.
3	12. "Reference drug product" shall mean the product to be manufactured
4	by the reference drug holder and includes both branded drugs of the NDA
5	holder and the biologic drug product of the biologic product license
б	applicant.
7	13. "Statutory exclusivity" shall mean those prohibitions on the
8	approval of drug applications under clauses (ii) through (iv) of section
9	505(c)(3)(E), section 527 or section 505A of the Federal Food, Drug, and
10	Cosmetic Act, 21 U.S.C. 355(c)(3)(E), on the licensing of biological
11	product applications under section 262(k)(7) of Title 42 of the United
12	States Code or section $262(m)(2)$ or (3) of Title 42 of the United States
13	Code.
14	§ 283. Preserving access to affordable drugs. 1. (a) Except as
15	provided in paragraph (c) of this subdivision, an agreement resolving or
16	settling, on a final or interim basis, a patent infringement claim, in
17	connection with the sale of a pharmaceutical product, shall be presumed
18	to have anticompetitive effects and shall be a violation of this section
19	if both of the following apply:
20	(i) A nonreference drug filer receives anything of value from another
21	company asserting patent infringement, including, but not limited to, an
22	exclusive license or a promise that the brand company will not launch an
23	authorized generic version of its brand drug; and
24	(ii) The nonreference drug filer agrees to limit or forego research,
25	development, manufacturing, marketing, or sales of the nonreference drug
26	filer's product for any period of time.
27	(b) As used in subparagraph (i) of paragraph (a) of this subdivision,
28	"anything of value" shall not include a settlement of a patent infringe-
29	ment claim in which the consideration granted by the brand or reference
30	drug filer to the nonreference drug filer as part of the resolution or
31	settlement consists of only one or more of the following:
32	(i) The right to market the competing product in the United States
33	before the expiration of either:
34	(A) A patent that is the basis for the patent infringement claim; or
35	(B) A patent right or other statutory exclusivity that would prevent
36	the marketing of the drug;
37	(ii) A covenant not to sue on a claim that the nonreference drug prod-
38	uct infringes a United States patent;
39	(iii) Compensation for saved reasonable future litigation expenses of
40	the reference drug holder but only if both of the following are true:
41	(A) The total compensation for saved litigation expenses is reflected
42	in budgets that the reference drug holder documented and adopted at
43	least six months before the settlement; and
44	(B) The compensation shall not exceed the lower of the following:
45	(1) Seven million five hundred thousand dollars; or
46	(2) Five percent of the revenue that the nonreference drug holder
47	projected or forecasted it would receive in the first three years of
48	sales of its version of the reference drug documented at least twelve
49	months before the settlement. If no projections or forecasts are avail-
50	able, the compensation shall not exceed two hundred fifty thousand
51	dollars;
52	(iv) An agreement resolving or settling a patent infringement claim
53	that permits a nonreference drug filer to begin selling, offering for
54	sale, or distributing the nonreference drug product if the reference
55	drug holder seeks approval to launch, obtains approval to launch, or
56	launches a different dosage, strength, or form of the reference drug

S. 4370

1	having the same active ingredient before the date set by the agreement
2	for entry of the nonreference drug filer. A different form of the refer-
3	ence drug shall not include an authorized generic version of the refer-
4	ence drug;
5	(v) An agreement by the reference drug holder not to interfere with
б	the nonreference drug filer's ability to secure and maintain regulatory
7	approval to market the nonreference drug product or an agreement to
8	facilitate the nonreference drug filer's ability to secure and maintain
9	regulatory approval to market the nonreference drug product; or
10	(vi) An agreement resolving a patent infringement claim in which the
11	reference drug holder forgives the potential damages accrued by a
12	nonreference drug holder for an at-risk launch of the nonreference drug
13	product that is the subject of that claim.
14	(c) Parties to an agreement are not in violation of paragraph (a) of
15	this subdivision if they can demonstrate by a preponderance of the
16	evidence that either of the following are met:
17	(i) The value received by the nonreference drug filer described in
18	subparagraph (i) of paragraph (a) of this subdivision is a fair and
19	reasonable compensation solely for other goods or services that the
20	nonreference drug filer has promised to provide; or
21	(ii) The agreement has directly generated procompetitive benefits and
22	the procompetitive benefits of the agreement outweigh the anticompet-
23	itive effects of the agreement.
24	2. In determining whether the parties to the agreement have met their
25	burden under paragraph (c) of subdivision one of this section, a court
26	of competent jurisdiction shall not presume any of the following:
27	(a) That entry into the marketplace could not have occurred until the
28	expiration of the relevant patent exclusivity or that the agreement's
29	provision for entry of the nonreference drug product before the expira-
30	tion of any patent exclusivity means that the agreement is procompet-
31	itive within the meaning of subparagraph (ii) of paragraph (c) of subdi-
32	vision one of this section;
33	(b) That any patent is enforceable and infringed by the nonreference
34	drug filer in the absence of a final adjudication binding on the filer
35	of those issues;
36	(c) That the agreement caused no delay in entry of the nonreference
37	drug filer's drug product because of the lack of Federal Food and Drug
38	Administration (FDA) approval of that or of another nonreference drug
39	product;
40	(d) That the agreement caused no harm or delay due to the possibility
41	that the nonreference drug filer's drug product might infringe some
42	patent that has not been asserted against the nonreference drug filer or
43	that is not subject to a final and binding adjudication on that filer as
44	to the patent's scope, enforceability, and infringement; or
45	(e) This subdivision shall not be construed to preclude a party from
46	introducing evidence regarding paragraphs (a), (b), (c) and (d) of this
47	subdivision, inclusive, and shall not be construed to preclude a court
48	of competent jurisdiction from making a determination regarding para-
49	graphs (a), (b), (c) and (d) of this subdivision, inclusive, based on
50	the full scope of the evidence.
51	3. In determining whether the parties to the agreement have met their
52	burden under paragraph (c) of subdivision one of this section, a court
53	of competent jurisdiction shall presume that the relevant product market
54	is that market consisting of the brand or reference drug of the company
55	alleging patent infringement and the drug product of the nonreference
56	company accused of infringement and any other biological product that is

1	licensed as biosimilar or is an AB-rated generic to the reference prod-
2	uct.
3	4. (a) This section shall not modify, impair, limit, or supersede the
4	applicability of the antitrust laws of the state pursuant to article
5	twenty-two of the general business law, unfair competition laws of the
6	state pursuant to article twenty-two-A of the general business law or
7	the availability of damages or remedies provided therein. This section
8	shall not modify, impair, limit, or supersede the right of any drug
9	company applicant to assert claims or counterclaims against any person,
10	under the antitrust laws or other laws relating to unfair competition of
11	<u>the federal antitrust law or state law.</u>
12	(b) If any provision of this subdivision, an amendment made to this
13	subdivision, or the application of any provision or amendment to any
14	person or circumstance is held to be unconstitutional, the remainder of
15	this subdivision, the amendments made to this subdivision, and the
16	application of the provisions of this subdivision or amendments to any
17	person or circumstance shall not be affected.
18	5. (a)(i) Each person that violates or assists in the violation of
19	this section shall forfeit and pay to the state a civil penalty suffi-
20	cient to deter violations of this section, as follows:
21	(A) If the person who violated this section received any value due to
22	that violation, an amount up to three times the value received by the
23	party that is reasonably attributable to the violation of this section,
24	or twenty million dollars, whichever is greater; or
25	(B) If the violator has not received anything of value as described in
26	this subparagraph, an amount up to three times the value given to other
27	parties to the agreement reasonably attributable to the violation of
28	this section, or twenty million dollars.
29	(C) For purposes of this subdivision, "reasonably attributable to the
30	violation" shall be determined by the state's share of the market for
31	the brand drug at issue in the agreement.
32	(ii) Any penalty described in subparagraph (i) of this paragraph shall
33	accrue only to the state and shall be recovered in a civil action
34	brought by the attorney general in its own name, or by any of its attor-
35	neys designated by it for that purpose, against any party to an agree-
36	ment that violates this section.
37	(b) Each party that violates or assists in the violation of this
38	section shall be liable for any damages, penalties, costs, fees, injunc-
39	tions, or other remedies that may be just and reasonable and available
40	under articles twenty-two or twenty-two-A of the general business law.
41	(c) If the state is awarded penalties under subparagraph (i) of para-
42	graph (a) of this subdivision, it shall not recover penalties pursuant
43	to another law identified in paragraph (b) of this subdivision. This
44	section shall not be construed to foreclose the state's ability to claim
45	any relief or damages available in paragraph (b) of this subdivision,
46	other than those that are penalties.
47	(d) An action to enforce a cause of action for a violation of this
48	section shall be commenced within four years after the cause of action
49	accrued.
50	§ 2. Severability clause. If any clause, sentence, paragraph, subdivi-
51	sion, section or part of this act shall be adjudged by any court of
52	competent jurisdiction to be invalid, such judgment shall not affect,
53	impair, or invalidate the remainder thereof, but shall be confined in
53 54	its operation to the clause, sentence, paragraph, subdivision, section
55	or part thereof directly involved in the controversy in which such judg-
56	ment shall have been rendered. It is hereby declared to be the intent of

1 the legislature that this act would have been enacted even if such 2 invalid provisions had not been included herein.

3 § 3. This act shall take effect on the one hundred eightieth day after 4 it shall have become a law.