

### 116TH CONGRESS 1ST SESSION

# H. R. 4398

To amend the Federal Trade Commission Act to prohibit anticompetitive behaviors by drug product manufacturers, and for other purposes.

### IN THE HOUSE OF REPRESENTATIVES

SEPTEMBER 19, 2019

Mr. Cicilline introduced the following bill; which was referred to the Committee on the Judiciary

# A BILL

To amend the Federal Trade Commission Act to prohibit anticompetitive behaviors by drug product manufacturers, and for other purposes.

- 1 Be it enacted by the Senate and House of Representa-
- 2 tives of the United States of America in Congress assembled,
- 3 SECTION 1. SHORT TITLE.
- 4 This Act may be cited as the "Affordable Prescrip-
- 5 tions for Patients Through Promoting Competition Act of
- 6 2019".
- 7 SEC. 2. PRODUCT HOPPING.
- 8 (a) In General.—The Federal Trade Commission
- 9 Act (15 U.S.C. 41 et seq.) is amended by inserting after
- 10 section 26 (15 U.S.C. 57c-2) the following:

## 1 "SEC. 27. PRODUCT HOPPING.

2	"(a) Definitions.—In this section:
3	"(1) Abbreviated New Drug application.—
4	The term 'abbreviated new drug application' means
5	an application under subsection (b)(2) or (j) of sec-
6	tion 505 of the Federal Food, Drug, and Cosmetic
7	Act (21 U.S.C. 355).
8	"(2) BIOSIMILAR BIOLOGICAL PRODUCT.—The
9	term 'biosimilar biological product' means a biologi-
10	cal product licensed under section 351(k) of the
11	Public Health Service Act (42 U.S.C. 262(k)).
12	"(3) BIOSIMILAR BIOLOGICAL PRODUCT LI-
13	CENSE APPLICATION.—The term 'biosimilar biologi-
14	cal product license application' means an application
15	submitted under section 351(k) of the Public Health
16	Service Act (42 U.S.C. 262(k)).
17	"(4) Follow-on product.—The term 'follow-
18	on product'—
19	"(A) means a drug approved through an
20	application or supplement to an application sub-
21	mitted under section 505(b) of the Federal
22	Food, Drug, and Cosmetic Act (21 U.S.C.
23	355(e)) or a biological product licensed through
24	an application or supplement to an application
25	submitted under section 351(a) of the Public
26	Health Service Act (42 U.S.C. 262(a)) for a

1	change, modification, or reformulation to the
2	same manufacturer's previously approved drug
3	or biological product that treats the same med-
4	ical condition; and
5	"(B) excludes such an application or sup-
6	plement to an application for a change, modi-
7	fication, or reformulation of a drug or biological
8	product that is requested by the Secretary or
9	necessary to comply with law, including sections
10	505A and 505B of the Federal Food, Drug,
11	and Cosmetic Act (21 U.S.C. 355a, 355c).
12	"(5) Generic drug.—The term 'generic drug'
13	means a drug approved under an application sub-
14	mitted under subsection (b)(2) or (j) of section $505$
15	of the Federal Food, Drug, and Cosmetic Act (21
16	U.S.C. 355).
17	"(6) Listed drug.—The term 'listed drug'
18	means a drug listed under section $505(j)(7)$ of the
19	Federal Food, Drug, and Cosmetic Act (21 U.S.C.
20	355(j)(7)).
21	"(7) Manufacturer.—The term 'manufac-
22	turer' means the holder, licensee, or assignee of—
23	"(A) an approved application for a drug
24	under section 505(c) of the Federal Food,
25	Drug, and Cosmetic Act (21 U.S.C. 355(c)); or

- 1 "(B) a biological product license under sec-2 tion 351(a) of the Public Health Service Act 3 (42 U.S.C. 262(a)).
  - "(8) REFERENCE PRODUCT.—The term 'reference product' has the meaning given the term in section 351(i) of the Public Health Service Act (42 U.S.C. 262(i)).
    - "(9) ULTIMATE PARENT ENTITY.—The term 'ultimate parent entity' has the meaning given the term in section 801.1 of title 16, Code of Federal Regulations, or any successor regulation.

### "(b) Prohibition on Product Hopping.—

"(1) PRIMA FACIE.—Except as provided in paragraph (2), a manufacturer of a reference product or listed drug shall be considered to have engaged in an unfair method of competition in or affecting commerce in violation of section 5(a) if the Commission demonstrates by a preponderance of the evidence in a proceeding initiated by the Commission under subsection (c)(1)(A), or in a suit brought under subparagraph (B) or (C) of subsection (c)(1), that, during the period beginning on the date on which the manufacturer of the reference product or listed drug first receives notice that an applicant has submitted to the Commissioner of Food and Drugs

1	an abbreviated new drug application or biosimilar bi-
2	ological product license application and ending on
3	the date that is 180 days after the date on which
4	that generic drug or biosimilar biological product is
5	first marketed, the manufacturer engaged in either
6	of the following actions:
7	"(A) The manufacturer engaged in a hard
8	switch, which shall be established by dem-
9	onstrating that the manufacturer engaged in ei-
10	ther of the following actions:
11	"(i) Upon the request of the manufac-
12	turer of the listed drug or reference prod-
13	uct, the Commissioner of Food and Drugs
14	withdrew the approval of the application
15	for the listed drug or reference product or
16	placed the listed drug or reference product
17	on the discontinued products list and the
18	manufacturer marketed or sold a follow-on
19	product.
20	"(ii)(I) The manufacturer of the listed
21	drug or reference product—
22	"(aa) announced withdrawal of,
23	discontinuance of the manufacture of,
24	or intent to withdraw the application
25	with respect to the drug or reference

1	product in a manner that impedes
2	competition from a generic drug or a
3	biosimilar biological product, as estab-
4	lished by objective circumstances; or
5	"(bb) destroyed the inventory of
6	the listed drug or reference product in
7	a manner that impedes competition
8	from a generic drug or a biosimilar bi-
9	ological product, which may be estab-
10	lished by objective circumstances; and
11	"(II) marketed or sold a follow-on
12	product.
13	"(B) The manufacturer engaged in a soft
14	switch, which shall be established by dem-
15	onstrating that the manufacturer engaged in
16	both of the following actions:
17	"(i) The manufacturer took actions
18	with respect to the listed drug or reference
19	product other than those described in sub-
20	paragraph (A) that unfairly disadvantage
21	the listed drug or reference product rel-
22	ative to the follow-on product described in
23	clause (ii) in a manner that impedes com-
24	petition from a generic drug or a bio-
25	similar biological product that is highly

1	similar to, and has no clinically meaningful
2	difference with respect to safety, purity,
3	and potency from, the reference product,
4	which may be established by objective cir-
5	cumstances.
6	"(ii) The manufacturer marketed or
7	sold a follow-on product.
8	"(2) Justification.—
9	"(A) In general.—Subject to paragraph
10	(3), the actions described in paragraph (1) by
11	a manufacturer of a listed drug or reference
12	product shall not be considered to be an unfair
13	method of competition in or affecting commerce
14	if—
15	"(i) the manufacturer demonstrates to
16	the Commission or a district court of the
17	United States, as applicable, by a prepon-
18	derance of the evidence in a proceeding ini-
19	tiated by the Commission under subsection
20	(c)(1)(A), or in a suit brought under sub-
21	paragraph (B) or (C) of subsection (c)(1),
22	that—
23	"(I) the manufacturer would
24	have taken the actions regardless of
25	whether a generic drug that ref-

1	erences the listed drug or biosimilar
2	biological product that references the
3	reference product had already entered
4	the market; and
5	"(II)(aa) with respect to a hard
6	switch under paragraph (1)(A), the
7	manufacturer took the action for rea-
8	sons relating to the safety risk to pa-
9	tients of the listed drug or reference
10	product;
11	"(bb) with respect to an action
12	described in item (aa) or (bb) of para-
13	graph (1)(A)(ii)(I), there is a supply
14	disruption that—
15	"(AA) is outside of the con-
16	trol of the manufacturer;
17	"(BB) prevents the produc-
18	tion or distribution of the appli-
19	cable listed drug or reference
20	product; and
21	"(CC) cannot be remedied
22	by reasonable efforts; or
23	"(cc) with respect to a soft
24	switch under paragraph (1)(B), the
25	manufacturer had legitimate pro-com-

1	petitive reasons, apart from the finan-
2	cial effects of reduced competition, to
3	take the action.
4	"(B) Rule of construction.—Nothing
5	in subparagraph (A) may be construed to limit
6	the information that the Commission may oth-
7	erwise obtain in any proceeding or action insti-
8	tuted with respect to a violation of this section.
9	"(3) Response.—With respect to a justifica-
10	tion offered by a manufacturer under paragraph (2),
11	the Commission may—
12	"(A) rebut any evidence presented by a
13	manufacturer during that justification; or
14	"(B) establish by a preponderance of the
15	evidence that, on balance, the pro-competitive
16	benefits from the conduct described in subpara-
17	graph (A) or (B) of paragraph (1), as applica-
18	ble, do not outweigh any anticompetitive effects
19	of the conduct, even in consideration of the jus-
20	tification so offered.
21	"(c) Enforcement.—
22	"(1) In general.—If the Commission has rea-
23	son to believe that any manufacturer has violated, is
24	violating, or is about to violate this section, the
25	Commission may take any of the following actions:

1	"(A) Institute a proceeding—
2	"(i) that, except as provided in para-
3	graph (2), complies with the requirements
4	under section 5(b); and
5	"(ii) in which the Commission may
6	impose on the manufacturer any penalty
7	that the Commission may impose for a vio-
8	lation of section 5.
9	"(B) In the same manner and to the same
10	extent as provided in section 13(b), bring suit
11	in a district court of the United States to tem-
12	porarily enjoin the action of the manufacturer.
13	"(C) Bring suit in a district court of the
14	United States, in which the Commission may
15	seek—
16	"(i) to permanently enjoin the action
17	of the manufacturer;
18	"(ii) any of the remedies described in
19	paragraph (3); and
20	"(iii) any other equitable remedy, in-
21	cluding ancillary equitable relief.
22	"(2) Judicial review.—
23	"(A) In general.—Notwithstanding any
24	provision of section 5, any manufacturer that is
25	subject to a final order of the Commission that

1	is issued in a proceeding initiated under para-
2	graph (1)(A) may, not later than 30 days after
3	the date on which the Commission issues the
4	order, petition for review of the order in—
5	"(i) the United States Court of Ap-
6	peals for the District of Columbia Circuit;
7	or
8	"(ii) the court of appeals of the
9	United States for the circuit in which the
10	ultimate parent entity of the manufacturer
11	is incorporated.
12	"(B) Treatment of findings.—In a re-
13	view of an order issued by the Commission con-
14	ducted by a court of appeals of the United
15	States under subparagraph (A), the factual
16	findings of the Commission shall be conclusive
17	if those facts are supported by the evidence.
18	"(3) Equitable remedies.—
19	"(A) DISGORGEMENT.—
20	"(i) In general.—In a suit brought
21	under paragraph (1)(C), the Commission
22	may seek, and the court may order,
23	disgorgement of any unjust enrichment
24	that a person obtained as a result of the
25	violation that gives rise to the suit.

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1	"(ii) Calculation.—Any disgorge-
2	ment that is ordered with respect to a per-
3	son under clause (i) shall be offset by any
4	amount of restitution ordered under sub-
5	paragraph (B).
6	"(iii) Limitations period.—The
7	Commission may seek disgorgement under
8	this subparagraph not later than 5 years
9	after the latest date on which the person
10	from which the disgorgement is sought re-
11	ceives any unjust enrichment from the ef-
12	fects of the violation that gives rise to the
13	suit in which the Commission seeks the
14	disgorgement.
15	"(B) Restitution.—
16	"(i) In general.—In a suit brought
17	under paragraph (1)(C), the Commission
18	may seek, and the court may order, res-
19	titution with respect to the violation that
20	gives rise to the suit.
21	"(ii) Limitations period.—The
22	Commission may seek restitution under
23	this subparagraph not later than 5 years
24	after the latest date on which the person

from which the restitution is sought re-

25

1	ceives any unjust enrichment from the ef-
2	fects of the violation that gives rise to the
3	suit in which the Commission seeks the
4	restitution.
5	"(4) Rules of Construction.—Nothing in
6	this subsection may be construed as—
7	"(A) requiring the Commission to bring a
8	suit seeking a temporary injunction under para-
9	graph (1)(B) before bringing a suit seeking a
10	permanent injunction under paragraph (1)(C);
11	or
12	"(B) affecting any other authority of the
13	Commission under this Act to seek relief or ob-
14	tain a remedy with respect to a violation of this
15	Act.".
16	(b) Applicability.—Section 27 of the Federal
17	Trade Commission Act, as added by subsection (a), shall
18	apply with respect to any—
19	(1) conduct that occurs on or after the date of
20	enactment of this Act; and
21	(2) action or proceeding that is commenced on
22	or after the date of enactment of this Act.
23	(c) Antitrust Laws.—Nothing in this section, or
24	the amendments made by this section, shall modify, im-
25	pair, limit, or supersede the applicability of the antitrust

- 1 laws as defined in subsection (a) of the first section of
- 2 the Clayton Act (15 U.S.C. 12(a)), and of section 5 of
- 3 the Federal Trade Commission Act (15 U.S.C. 45) to the
- 4 extent that it applies to unfair methods of competition.
- 5 (d) Rulemaking.—The Federal Trade Commission
- 6 may issue rules under section 553 of title 5, United States
- 7 Code, to carry out section 27 of the Federal Trade Com-
- 8 mission Act, as added by subsection (a), including by de-
- 9 fining any terms used in such section 27 (other than terms
- 10 that are defined in subsection (a) of such section 27).

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