

115TH CONGRESS 1ST SESSION

H. R. 2212

To promote competition in the market for drugs and biological products by facilitating the timely entry of lower-cost generic and biosimilar versions of those drugs and biological products.

IN THE HOUSE OF REPRESENTATIVES

April 27, 2017

Mr. Marino (for himself and Mr. Cicilline) introduced the following bill; which was referred to the Committee on Energy and Commerce, and in addition to the Committee on the Judiciary, for a period to be subsequently determined by the Speaker, in each case for consideration of such provisions as fall within the jurisdiction of the committee concerned

A BILL

To promote competition in the market for drugs and biological products by facilitating the timely entry of lowercost generic and biosimilar versions of those drugs and biological products.

- 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 "Creating and Restoring
- 5 Equal Access To Equivalent Samples Act of 2017" or the
- 6 "CREATES Act of 2017".

1 SEC. 2. FINDINGS.

- 2 Congress finds the following:
- (1) It is the policy of the United States to promote competition in the market for drugs and biological products by facilitating the timely entry of low-cost generic and biosimilar versions of those drugs and biological products.
 - (2) Since their enactment in 1984 and 2010, respectively, the Drug Price Competition and Patent Term Restoration Act of 1984 (Public Law 98–417; 98 Stat. 1585) and the Biologics Price Competition and Innovation Act of 2009 (Subtitle A of title VII of Public Law 111–148; 124 Stat. 804), have provided pathways for making lower-cost versions of previously approved drugs and previously licensed biological products available to the people of the United States in a timely manner, thereby lowering overall prescription drug costs for patients and tax-payers by billions of dollars each year.
 - (3) In order for these pathways to function as intended, developers of generic drugs and biosimilar biological products (referred to in this section as "generic product developers") must be able to obtain quantities of the reference listed drug or biological product with which the generic drug or biosimilar biological product is intended to compete (referred to

- in this section as a "covered product") for purposes of supporting an application for approval by the Food and Drug Administration, including for testing to show that—
 - (A) a prospective generic drug is bioequivalent to the covered product in accordance with subsection (j) of section 505 of the Federal, Food, Drug, and Cosmetic Act (21 U.S.C. 355), or meets the requirements for approval of an application submitted under subsection (b)(2) of that section; or
 - (B) a prospective biosimilar biological product is biosimilar to or interchangeable with its reference biological product under section 351(k) of the Public Health Service Act (42 U.S.C. 262(k)), as applicable.
 - (4) For drugs and biological products that are subject to a risk evaluation and mitigation strategy, another essential component in the creation of low-cost generic and biosimilar versions of covered products is the ability of generic product developers to join the manufacturer of the covered product (referred to in this section as the "license holder") in a single, shared system of elements to assure safe use and supporting agreements, or secure a variance

- 1 therefrom, as required by section 505–1 of the Fed-2 eral Food, Drug, and Cosmetic Act (21 U.S.C. 355– 1). 3
- (5) Contrary to the policy of the United States 5 to promote competition in the market for drugs and 6 biological products by facilitating the timely entry of 7 lower-cost generic and biosimilar versions of those 8 drugs and biological products, certain license holders 9 are preventing generic product developers from ob-10 taining quantities of the covered product necessary for the generic product developer to support an ap-12 plication for approval by the Food and Drug Admin-13 istration, including testing to show bioequivalence, 14 biosimilarity, or interchangeability to the covered 15 product, in some instances based on the justification 16 that the covered product is subject to a risk evalua-17 tion and mitigation strategy with elements to assure 18 safe use under section 505–1 of the Federal Food, 19 Drug, and Cosmetic Act (21 U.S.C. 355–1).
 - (6) The Director of the Center for Drug Evaluation and Research at the Food and Drug Administration has testified that some manufacturers of covered products have used REMS and distribution restrictions adopted by the manufacturer on their own behalf as reasons to not sell quantities of a cov-

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- ered product to generic product developers, causing barriers and delays in getting generic products on the market. The Food and Drug Administration has reported receiving significant numbers of inquiries from generic product developers who were unable to obtain samples of covered products to conduct necessary testing and otherwise meet requirements for approval of generic drugs.
 - (7) The Chairwoman of the Federal Trade Commission has testified that the Federal Trade Commission continues to be very concerned about potential abuses by manufacturers of brand drugs of REMS or other closed distribution systems to impede generic competition.
 - (8) Also contrary to the policy of the United States to promote competition in the market for drugs and biological products by facilitating the timely entry of lower-cost generic and biosimilar versions of those drugs and biological products, certain license holders are impeding the prompt negotiation and development on commercially reasonable terms of a single, shared system of elements to assure safe use, which may be necessary for the generic product developer to gain approval for its drug or licensing for its biological product.

(9) While the antitrust laws may address the refusal by some license holders to provide quantities of a covered product to a generic product developer, a more tailored legal pathway would help ensure that generic product developers can obtain necessary quantities of a covered product in a timely way for purposes of developing a generic drug or biosimilar biological product, facilitating competition in the marketplace for drugs and biological products.

(10) The antitrust laws may address actions by license holders who impede the prompt negotiation and development of a single, shared system of elements to assure safe use, and the Food and Drug Administration has some authority to waive the requirement of a single, shared system. Clearer regulatory authority to approve different systems that meet the statutory requirements to ensure patient safety, however, would limit the effectiveness of bad faith negotiations over single, shared systems to delay generic approval. At the same time, clearer regulatory authority would ensure all systems protect patient safety.

23 SEC. 3. ACTIONS FOR DELAYS OF GENERIC DRUGS AND

24 BIOSIMILAR BIOLOGICAL PRODUCTS.

25 (a) Definitions.—In this section—

1	(1) the term "covered product"—
2	(A) means—
3	(i) any drug approved under sub-
4	section (b) or (j) of section 505 of the Fed-
5	eral Food, Drug, and Cosmetic Act (21
6	U.S.C. 355) or biological product licensed
7	under subsection (a) or (k) of section 351
8	of the Public Health Service Act (42
9	U.S.C. 262);
10	(ii) any combination of a drug or bio-
11	logical product described in clause (i); or
12	(iii) when reasonably necessary to
13	demonstrate sameness, biosimilarity, or
14	interchangeability for purposes of section
15	505 of the Federal Food, Drug, and Cos-
16	metic Act (21 U.S.C. 355), or section 351
17	of the Public Health Service Act (42
18	U.S.C. 262), as applicable, any product,
19	including any device, that is marketed or
20	intended for use with such drug or biologi-
21	cal product; and
22	(B) does not include any drug or biological
23	product that the Secretary has determined to be
24	currently in shortage and that appears on the
25	drug shortage list in effect under section 506E

1	of the Federal Food, Drug, and Cosmetic Act
2	(21 U.S.C. 356e), unless the shortage will not
3	be promptly resolved—
4	(i) as demonstrated by the fact that
5	the drug or biological product has been in
6	shortage for more than 6 months; or
7	(ii) as otherwise determined by the
8	Secretary;
9	(2) the term "device" has the meaning given
10	the term in section 201 of the Federal Food, Drug,
11	and Cosmetic Act (21 U.S.C. 321);
12	(3) the term "eligible product developer" means
13	a person that seeks to develop a product for ap-
14	proval pursuant to an application for approval under
15	subsection (b)(2) or (j) of section 505 of the Federal
16	Food, Drug, and Cosmetic Act (21 U.S.C. 355) or
17	for licensing pursuant to an application under sec-
18	tion 351(k) of the Public Health Service Act (42
19	U.S.C. 262(k));
20	(4) the term "license holder" means the holder
21	of an application approved under subsection (c) or
22	(j) of section 505 of the Federal Food, Drug, and
23	Cosmetic Act (21 U.S.C. 355) or the holder of a li-
24	cense under subsection (a) or (k) of section 351 of

1	the Public Health Service Act (42 U.S.C. 262) for
2	a covered product;
3	(5) the term "REMS" means a risk evaluation
4	and mitigation strategy under section 505-1 of the
5	Federal Food, Drug, and Cosmetic Act (21 U.S.C.
6	355–1);
7	(6) the term "REMS with ETASU" means a
8	REMS that contains elements to assure safe use
9	under section 505–1 of the Federal Food, Drug, and
10	Cosmetic Act (21 U.S.C. 355–1);
11	(7) the term "Secretary" means the Secretary
12	of Health and Human Services;
13	(8) the term "single, shared system of elements
14	to assure safe use" means a single, shared system
15	of elements to assure safe use under section 505–1
16	of the Federal Food, Drug, and Cosmetic Act (21
17	U.S.C. 355–1); and
18	(9) the term "sufficient quantities" means an
19	amount of a covered product that allows the eligible
20	product developer to—
21	(A) conduct testing to support an applica-
22	tion—
23	(i) for approval under subsection
24	(b)(2) or (i) of section 505 of the Federal

1	Food, Drug, and Cosmetic Act (21 U.S.C
2	355); or
3	(ii) for licensing under section 351(k)
4	of the Public Health Service Act (42
5	U.S.C. 262(k)); and
6	(B) fulfill any regulatory requirements re-
7	lating to such an application for approval or li-
8	censing.
9	(b) Civil Action for Failure To Provide Suffi-
10	CIENT QUANTITIES OF A COVERED PRODUCT.—
11	(1) IN GENERAL.—An eligible product developer
12	may bring a civil action against the license holder
13	for a covered product seeking relief under this sub-
14	section in an appropriate district court of the United
15	States alleging that the license holder has declined
16	to provide sufficient quantities of the covered prod-
17	uct to the eligible product developer on commercially
18	reasonable, market-based terms.
19	(2) Elements.—
20	(A) In general.—To prevail in a civil ac-
21	tion brought under paragraph (1), an eligible
22	product developer shall prove, by a preponder-
23	ance of the evidence—
24	(i) that—

1	(I) the covered product is not
2	subject to a REMS with ETASU; or
3	(II) if the covered product is sub-
4	ject to a REMS with ETASU—
5	(aa) the eligible product de-
6	veloper has obtained a covered
7	product authorization from the
8	Secretary in accordance with sub-
9	paragraph (B); and
10	(bb) the eligible product de-
11	veloper has provided a copy of
12	the covered product authorization
13	to the license holder;
14	(ii) that, as of the date on which the
15	civil action is filed, the product developer
16	has not obtained sufficient quantities of
17	the covered product on commercially rea-
18	sonable, market-based terms;
19	(iii) that the eligible product developer
20	has requested to purchase sufficient quan-
21	tities of the covered product from the li-
22	cense holder; and
23	(iv) that the license holder has not de-
24	livered to the eligible product developer
25	sufficient quantities of the covered product

1	on commercially reasonable, market-based
2	terms—
3	(I) for a covered product that is
4	not subject to a REMS with ETASU,
5	by the date that is 31 days after the
6	date on which the license holder re-
7	ceived the request for the covered
8	product; and
9	(II) for a covered product that is
10	subject to a REMS with ETASU, by
11	31 days after the later of—
12	(aa) the date on which the
13	license holder received the re-
14	quest for the covered product; or
15	(bb) the date on which the
16	license holder received a copy of
17	the covered product authorization
18	issued by the Secretary in ac-
19	cordance with subparagraph (B).
20	(B) Authorization for covered prod-
21	UCT SUBJECT TO A REMS WITH ETASU.—
22	(i) Request.—An eligible product de-
23	veloper may submit to the Secretary a
24	written request for the eligible product de-
25	veloper to be authorized to obtain suffi-

1	cient quantities of an individual covered
2	product subject to a REMS with ETASU.
3	(ii) Authorization.—Not later than
4	90 days after the date on which a request
5	under clause (i) is received, the Secretary
6	shall, by written notice, authorize the eligi-
7	ble product developer to obtain sufficient
8	quantities of an individual covered product
9	subject to a REMS with ETASU for pur-
10	poses of—
11	(I) development and testing that
12	does not involve human clinical trials,
13	if the eligible product developer has
14	agreed to comply with any conditions
15	the Secretary determines necessary; or
16	(II) development and testing that
17	involves human clinical trials, if the
18	eligible product developer has—
19	(aa)(AA) submitted proto-
20	cols, informed consent docu-
21	ments, and informational mate-
22	rials for testing that include pro-
23	tections that provide safety pro-
24	tections comparable to those pro-

1	vided by the REMS for the cov-
2	ered product; or
3	(BB) otherwise satisfied the
4	Secretary that such protections
5	will be provided; and
6	(bb) met any other require-
7	ments the Secretary may estab-
8	lish.
9	(iii) Notice.—A covered product au-
10	thorization issued under this subparagraph
11	shall state that the provision of the covered
12	product by the license holder under the
13	terms of the authorization will not be a
14	violation of the REMS for the covered
15	product.
16	(3) Affirmative Defense.—In a civil action
17	brought under paragraph (1), it shall be an affirma-
18	tive defense, on which the defendant has the burden
19	of persuasion by a preponderance of the evidence—
20	(A) that, on the date on which the eligible
21	product developer requested to purchase suffi-
22	cient quantities of the covered product from the
23	license holder—
24	(i) neither the license holder nor any
25	of its agents, wholesalers, or distributors

1	was engaged in the manufacturing or com-
2	mercial marketing of the covered product
3	and
4	(ii) neither the license holder nor any
5	of its agents, wholesalers, or distributors
6	otherwise had access to inventory of the
7	covered product to supply to the eligible
8	product developer on commercially reason-
9	able, market-based terms; or
10	(B) that—
11	(i) the license holder sells the covered
12	product through agents, distributors, or
13	wholesalers;
14	(ii) the license holder has placed no
15	restrictions, explicit or implicit, on its
16	agents, distributors, or wholesalers to sell
17	covered products to eligible product devel-
18	opers; and
19	(iii) the covered product can be pur-
20	chased by the eligible product developer in
21	sufficient quantities on commercially rea-
22	sonable, market-based terms from the
23	agents, distributors, or wholesalers of the
24	license holder.
25	(4) Remedies.—

1	(A) In General.—If an eligible product
2	developer prevails in a civil action brought
3	under paragraph (1), the court shall—
4	(i) order the license holder to provide
5	to the eligible product developer without
6	delay sufficient quantities of the covered
7	product on commercially reasonable, mar-
8	ket-based terms;
9	(ii) award to the eligible product de-
10	veloper reasonable attorney fees and costs
11	of the civil action; and
12	(iii) award to the eligible product de-
13	veloper a monetary amount sufficient to
14	deter the license holder from failing to pro-
15	vide other eligible product developers with
16	sufficient quantities of a covered product
17	on commercially reasonable, market-based
18	terms, if the court finds, by a preponder-
19	ance of the evidence—
20	(I) that the license holder delayed
21	providing sufficient quantities of the
22	covered product to the eligible product
23	developer without a legitimate busi-
24	ness justification; or

1	(II) that the license holder failed
2	to comply with an order issued under
3	clause (i).
4	(B) MAXIMUM MONETARY AMOUNT.—A
5	monetary amount awarded under subparagraph
6	(A)(iii) shall not be greater than the revenue
7	that the license holder earned on the covered
8	product during the period—
9	(i) beginning on—
10	(I) for a covered product that is
11	not subject to a REMS with ETASU,
12	the date that is 31 days after the date
13	on which the license holder received
14	the request; or
15	(II) for a covered product that is
16	subject to a REMS with ETASU, the
17	date that is 31 days after the later
18	of—
19	(aa) the date on which the
20	license holder received the re-
21	quest; or
22	(bb) the date on which the
23	license holder received a copy of
24	the covered product authorization
25	issued by the Secretary in ac-

1	cordance with paragraph (2)(B);
2	and
3	(ii) ending on the date on which the
4	eligible product developer received suffi-
5	cient quantities of the covered product.
6	(C) AVOIDANCE OF DELAY.—The court
7	may issue an order under subparagraph (A)(i)
8	before conducting further proceedings that may
9	be necessary to determine whether the eligible
10	product developer is entitled to an award under
11	clause (ii) or (iii) of subparagraph (A), or the
12	amount of any such award.
13	(c) Limitation of Liability.—A license holder for
14	a covered product shall not be liable for any claim arising
15	out of the failure of an eligible product developer to follow
16	adequate safeguards to assure safe use of the covered
17	product during development or testing activities described
18	in this section, including transportation, handling, use, or
19	disposal of the covered product by the eligible product de-
20	veloper.
21	(d) Rule of Construction.—
22	(1) Definition.—In this subsection, the term
23	"antitrust laws"—

1	(A) has the meaning given the term in
2	subsection (a) of the first section of the Clayton
3	Act (15 U.S.C. 12); and
4	(B) includes section 5 of the Federal
5	Trade Commission Act (15 U.S.C. 45) to the
6	extent that such section applies to unfair meth-
7	ods of competition.
8	(2) Antitrust laws.—Nothing in this section
9	shall be construed to limit the operation of any pro-
10	vision of the antitrust laws.
11	SEC. 4. REMS APPROVAL PROCESS FOR SUBSEQUENT FIL-
12	ERS.
13	Section 505–1 of the Federal Food, Drug, and Cos-
14	metic Act (21 U.S.C. 355–1) is amended—
15	(1) in subsection $(g)(4)(B)$ —
16	(A) in clause (i) by striking "or" after the
17	semicolon;
18	(B) in clause (ii) by striking the period at
19	the end and inserting "; or"; and
20	(C) by adding at the end the following:
21	"(iii) accommodate different approved
22	risk evaluation and mitigation strategies
23	for a reference drug product and a drug
24	that is the subject of an abbreviated new
25	drug application."; and

1	(2) in subsection (i)(1), by striking subpara-
2	graph (B) and inserting the following:
3	"(B) Elements to assure safe use, if re-
4	quired under subsection (f) for the listed drug.
5	"(i) Subject to clause (ii), a drug that
6	is the subject of an abbreviated new drug
7	application may use—
8	"(I) a single, shared system with
9	the listed drug under subsection (f);
10	or
11	"(II) a different, comparable as-
12	pect of the elements to assure safe use
13	under subsection (f).
14	"(ii) The Secretary may require a
15	drug that is the subject of an abbreviated
16	new drug application and the listed drug to
17	use a single, shared system under sub-
18	section (f), if the Secretary determines
19	that no different, comparable aspect of the
20	elements to assure safe use could satisfy
21	the requirements of subsection (f).".