

115TH CONGRESS 2D SESSION

S. 2476

To amend the Federal Food, Drug, and Cosmetic Act to ensure that valid generic drugs may enter the market.

IN THE SENATE OF THE UNITED STATES

February 28, 2018

Ms. Smith introduced the following bill; which was read twice and referred to the Committee on Health, Education, Labor, and Pensions

A BILL

To amend the Federal Food, Drug, and Cosmetic Act to ensure that valid generic drugs may enter the market.

- 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 "Expanding Access to
- 5 Low Cost Generic Drugs Act".
- 6 SEC. 2. 180-DAY EXCLUSIVITY PERIOD AMENDMENTS RE-
- 7 GARDING FIRST APPLICANT STATUS.
- 8 (a) Amendments to Federal Food, Drug, and
- 9 Cosmetic Act.—

1	(1) In General.—Section $505(j)(5)(B)$ of the
2	Federal Food, Drug, and Cosmetic Act (21 U.S.C.
3	355(j)(5)(B)) is amended—
4	(A) in clause (iv)(II)—
5	(i) by striking item (bb); and
6	(ii) by redesignating items (cc) and
7	(dd) as items (bb) and (cc), respectively;
8	(B) by redesignating clause (v) as clause
9	(ix); and
10	(C) by inserting after clause (iv) the fol-
11	lowing:
12	"(v) First applicant defined.—As used in
13	this subsection, the term 'first applicant' means an
14	applicant—
15	"(I)(aa) that, on the first day on which a
16	substantially complete application containing a
17	certification described in paragraph
18	(2)(A)(vii)(IV) is submitted for approval of a
19	drug, submits a substantially complete applica-
20	tion that contains and lawfully maintains a cer-
21	tification described in paragraph (2)(A)(vii)(IV)
22	for the drug; and
23	"(bb) that has not entered into a disquali-
24	fying agreement described under clause
25	(vii)(II); or

1	"(II)(aa) for the drug that is not described
2	in subclause (I) and that, with respect to the
3	applicant and drug, each requirement described
4	in clause (vi) is satisfied; and
5	"(bb) that has not entered into a disquali-
6	fying agreement described under clause
7	(vii)(II).
8	"(vi) Requirements de-
9	scribed in this clause are the following:
10	"(I) The applicant described in clause
11	(v)(II) submitted and lawfully maintains a cer-
12	tification described in paragraph (2)(A)(vii)(IV)
13	or a statement described in paragraph
14	(2)(A)(viii) for each unexpired patent for which
15	a first applicant described in clause (v)(I) had
16	submitted a certification described in paragraph
17	(2)(A)(vii)(IV) on the first day on which a sub-
18	stantially complete application containing such
19	a certification was submitted.
20	"(II) With regard to each such unexpired
21	patent for which the applicant described in
22	clause (v)(II) submitted a certification de-
23	scribed in paragraph (2)(A)(vii)(IV), no action
24	for patent infringement was brought against

such applicant within the 45-day period speci-

1 fied in clause (iii); or if an action was brought 2 within such time period, such an action was 3 withdrawn or dismissed by a court (including a 4 district court) without a decision that the patent was valid and infringed; or if an action was 6 brought within such time period and was not 7 withdrawn or so dismissed, such applicant has 8 obtained the decision of a court (including a 9 district court) that the patent is invalid or not infringed (including any substantive determina-10 tion that there is no cause of action for patent 12 infringement or invalidity, and including a set-13 tlement order or consent decree signed and en-14 tered by the court stating that the patent is in-15 valid or not infringed).

> "(III) If an applicant described in clause (v)(I) has begun commercial marketing of such drug, the applicant described in clause (v)(II) does not begin commercial marketing of such drug until the date that is 30 days after the date on which the applicant described in clause (v)(I) began such commercial marketing.".

(2)Conforming AMENDMENT.—Section 505(j)(5)(D)(i)(IV)Act (21of such U.S.C. 355(j)(5)(D)(i)(IV) is amended by striking "The

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1	first applicant" and inserting "The first applicant
2	as defined in subparagraph (B)(v)(I),".
3	(b) APPLICABILITY.—The amendments made by sub-
4	section (a) shall apply only with respect to an application
5	filed under section 505(j) of the Federal Food, Drug, and
6	Cosmetic Act (21 U.S.C. 355(j)) to which the amendments
7	made by section 1102(a) of the Medicare Prescription
8	Drug, Improvement, and Modernization Act of 2003 (Pub-
9	lic Law 108–173) apply.
10	SEC. 3. 180-DAY EXCLUSIVITY PERIOD AMENDMENTS RE-
11	GARDING AGREEMENTS TO DEFER COMMER
12	CIAL MARKETING.
13	(a) Amendments to Federal Food, Drug, and
13 14	(a) Amendments to Federal Food, Drug, and Cosmetic Act.—
14	COSMETIC ACT.—
14 15	Cosmetic Act.— (1) Limitations on agreements to defer
14 15 16	COSMETIC ACT.— (1) LIMITATIONS ON AGREEMENTS TO DEFER COMMERCIAL MARKETING DATE.—Section
14 15 16 17	COSMETIC ACT.— (1) LIMITATIONS ON AGREEMENTS TO DEFER COMMERCIAL MARKETING DATE.—Section 505(j)(5)(B) of the Federal Food, Drug, and Cosmercial Food, Drug, Drug, Cosmercial Food, Drug, Cosme
14 15 16 17	COSMETIC ACT.— (1) LIMITATIONS ON AGREEMENTS TO DEFER COMMERCIAL MARKETING DATE.—Section 505(j)(5)(B) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 355(j)(5)(B)), as amended by
114 115 116 117 118	(1) Limitations on agreements to defer commercial marketing date.—Section 505(j)(5)(B) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 355(j)(5)(B)), as amended by section 2, is further amended by adding at the end
14 15 16 17 18 19 20	(1) Limitations on agreements to defer commercial marketing date.—Section 505(j)(5)(B) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 355(j)(5)(B)), as amended by section 2, is further amended by adding at the end the following:
14 15 16 17 18 19 20 21	(1) Limitations on agreements to defer commercial marketing date.—Section 505(j)(5)(B) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 355(j)(5)(B)), as amended by section 2, is further amended by adding at the end the following: "(vii) Agreement by first applicant to

1 "(I) AGREEMENT TO DEFER APPROVAL OR 2 COMMERCIAL MARKETING DATE.—An agree-3 ment described in this subclause is an agree-4 ment between a first applicant and the holder 5 of the application for the listed drug or an 6 owner of one or more of the patents as to which 7 any applicant submitted a certification quali-8 fying such applicant for the 180-day exclusivity 9 period whereby that applicant agrees, directly 10 or indirectly, (aa) not to seek an approval of its 11 application that is made effective on the earliest 12 possible date under this subparagraph, subpara-13 graph (F) of this paragraph, section 505A, or 14 section 527, (bb) not to begin the commercial 15 marketing of its drug on the earliest possible 16 date after receiving an approval of its applica-17 tion that is made effective under this subpara-18 graph, subparagraph (F) of this paragraph, sec-19 tion 505A, or section 527, or (cc) to both items 20 (aa) and (bb). 21 "(II) AGREEMENT THAT DISQUALIFIES AP-22

"(II) AGREEMENT THAT DISQUALIFIES AP-PLICANT FROM FIRST APPLICANT STATUS.—An agreement described in this subclause is an agreement between an applicant and the holder of the application for the listed drug or an

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1 owner of one or more of the patents as to which 2 any applicant submitted a certification quali-3 fying such applicant for the 180-day exclusivity 4 period whereby that applicant agrees, directly 5 or indirectly, not to seek an approval of its ap-6 plication or not to begin the commercial marketing of its drug until a date that is after the 7 8 expiration of the 180-day exclusivity period 9 awarded to another applicant with respect to 10 such drug (without regard to whether such 180-11 day exclusivity period is awarded before or after 12 the date of the agreement). "(viii) Limitation on acceleration.—If an 13 agreement described in clause (vii)(I) includes more 14 15 than 1 possible date when an applicant may seek an 16

approval of its application or begin the commercial marketing of its drug—

"(I) the applicant may seek an approval of its application or begin such commercial marketing on the date that is the earlier of—

> "(aa) the latest date set forth in the agreement on which that applicant can receive an approval that is made effective under this subparagraph, subparagraph (F) of this paragraph, section 505A, or

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1	section 527, or begin the commercial mar-
2	keting of such drug, without regard to any
3	other provision of such agreement pursu-
4	ant to which the commercial marketing
5	could begin on an earlier date; or
6	"(bb) 180 days after another first ap-
7	plicant begins commercial marketing of
8	such drug; and
9	"(II) the latest date set forth in the agree-
10	ment on which that applicant can receive an ap-
11	proval that is made effective under this sub-
12	paragraph, subparagraph (F) of this paragraph
13	section 505A, or section 527, or begin the com-
14	mercial marketing of such drug, without regard
15	to any other provision of such agreement pursu-
16	ant to which commercial marketing could begin
17	on an earlier date, shall be the date used to de-
18	termine whether an applicant is disqualified
19	from first applicant status pursuant to clause
20	(vii)(II).''.
21	(2) Notification of fda.—Section 505(j) of
22	such Act (21 U.S.C. 355(j)) is amended by adding
23	at the end the following:

- 1 "(14)(A) The holder of an abbreviated application
- 2 under this subsection shall submit to the Secretary a noti-
- 3 fication that includes—
- 4 "(i)(I) the text of any agreement entered into
- 5 by such holder described under paragraph
- 6 (5)(B)(vii)(I); or
- 7 "(II) if such an agreement has not been re-
- 8 duced to text, a written detailed description of such
- 9 agreement that is sufficient to disclose all the terms
- and conditions of the agreement; and
- 11 "(ii) the text, or a written detailed description
- in the event of an agreement that has not been re-
- duced to text, of any other agreements that are con-
- tingent upon, provide a contingent condition for, or
- are otherwise related to an agreement described in
- 16 clause (i).
- 17 "(B) The notification described under subparagraph
- 18 (A) shall be submitted not later than 10 business days
- 19 after execution of the agreement described in subpara-
- 20 graph (A)(i). Such notification is in addition to any notifi-
- 21 cation required under section 1112 of the Medicare Pre-
- 22 scription Drug, Improvement, and Modernization Act of
- 23 2003.
- 24 "(C) Any information or documentary material filed
- 25 with the Secretary pursuant to this paragraph shall be ex-

- 1 empt from disclosure under section 552 of title 5, United
- 2 States Code, and no such information or documentary ma-
- 3 terial may be made public, except as may be relevant to
- 4 any administrative or judicial action or proceeding. Noth-
- 5 ing in this paragraph is intended to prevent disclosure to
- 6 either body of the Congress or to any duly authorized com-
- 7 mittee or subcommittee of the Congress.".
- 8 (3) Prohibited acts.—Section 301(e) of such
- 9 Act (21 U.S.C. 331(e)) is amended by striking "505
- 10 (i) or (k)" and inserting "505 (i), (j)(14), or (k)".
- 11 (b) Infringement of Patent.—Section 271(e) of
- 12 title 35, United States Code, is amended by adding at the
- 13 end the following:
- 14 "(7) The exclusive remedy under this section for an
- 15 infringement of a patent for which the Secretary of Health
- 16 and Human Services has published information pursuant
- 17 to subsection (b)(1) or (c)(2) of section 505 of the Federal
- 18 Food, Drug, and Cosmetic Act shall be an action brought
- 19 under this subsection within the 45-day period described
- 20 in subsection (j)(5)(B)(iii) or (e)(3)(C) of section 505 of
- 21 the Federal Food, Drug, and Cosmetic Act.".
- 22 (c) Applicability.—
- 23 (1) Limitations on acceleration of De-
- 24 FERRED COMMERCIAL MARKETING DATE.—The

1	amendment made by subsection (a)(1) shall apply
2	only with respect to—

- (A) an application filed under section 505(j) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 355(j)) to which the amendments made by section 1102(a) of the Medicare Prescription Drug, Improvement, and Modernization Act of 2003 (Public Law 108–173) apply; and
- (B) an agreement described under section 505(j)(5)(B)(vii)(I) of the Federal Food, Drug, and Cosmetic Act (as added by subsection (a)(1)) executed after the date of enactment of this Act.
- (2) Notification of fda.—The amendments made by paragraphs (2) and (3) of subsection (a) shall apply only with respect to an agreement described under section 505(j)(5)(B)(vii)(I) of the Federal Food, Drug, and Cosmetic Act (as added by subsection (a)(1)) executed after the date of enactment of this Act.

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