

115TH CONGRESS
2D SESSION

H. R. 5811

To amend the Federal Food, Drug, and Cosmetic Act with respect to postapproval study requirements for certain controlled substances, and for other purposes.

IN THE HOUSE OF REPRESENTATIVES

MAY 15, 2018

Mr. MCNERNEY (for himself and Mr. GRIFFITH) introduced the following bill;
which was referred to the Committee on Energy and Commerce

A BILL

To amend the Federal Food, Drug, and Cosmetic Act with respect to postapproval study requirements for certain controlled substances, 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 “Long-Term Opioid Ef-
5 ficacy Act of 2018”.

6 **SEC. 2. POSTAPPROVAL STUDY REQUIREMENTS.**

7 (a) PURPOSES OF STUDY.—Section 505(o)(3)(B) of
8 the Federal Food, Drug, and Cosmetic Act (21 U.S.C.

1 355(o)(3)(B)) is amended by adding at the end the fol-
2 lowing:

3 “(iv) To assess a potential reduction
4 in effectiveness or an increase in serious
5 risk of the drug for the conditions of use
6 prescribed, recommended, or suggested in
7 the labeling thereof if—

8 “(I) the drug involved—

9 “(aa) is or contains a sub-
10 stance for which a listing in any
11 schedule is in effect (on a tem-
12 porary or permanent basis) under
13 section 201 of the Controlled
14 Substances Act; or

15 “(bb) is a drug that has not
16 been approved under this section
17 or licensed under section 351 of
18 the Public Health Service Act,
19 for which an application for such
20 approval or licensure is pending
21 or anticipated, and for which the
22 Secretary provides notice to the
23 sponsor that the Secretary in-
24 tends to issue a scientific and
25 medical evaluation and rec-

1 commend controls under the Con-
2 trolled Substances Act; and
3 “(II) the potential reduction in
4 effectiveness or the increase in serious
5 risk could result in the benefits of the
6 drug no longer outweighing the
7 risks.”.

8 (b) ESTABLISHMENT OF REQUIREMENT.—Section
9 505(o)(3)(C) of the Federal Food, Drug, and Cosmetic
10 Act (21 U.S.C. 355(o)(3)(C)) is amended by striking
11 “such requirement” and all that follows through “safety
12 information.” and inserting the following: “such require-
13 ment—

14 “(i) in the case of a purpose described
15 in clause (i), (ii), or (iii) of subparagraph
16 (B), only if the Secretary becomes aware of
17 new safety information; and

18 “(ii) in the case of a purpose de-
19 scribed in clause (iv) of such subpara-
20 graph, if the Secretary determines that
21 new effectiveness information exists.”.

22 (c) APPLICABILITY.—Section 505(o)(3) of the Fed-
23 eral Food, Drug, and Cosmetic Act (21 U.S.C. 355(o)(3))
24 is amended by adding at the end the following new sub-
25 paragraph:

1 “(E) APPLICABILITY.—The conduct of a
2 study or clinical trial required pursuant to this
3 paragraph for the purpose specified in subpara-
4 graph (B)(iv) shall not be considered a new
5 clinical investigation for the purpose of a period
6 of exclusivity under clause (iii) or (iv) of sub-
7 section (c)(3)(E) or clause (iii) or (iv) of sub-
8 section (j)(5)(F).”.

9 (d) NEW EFFECTIVENESS INFORMATION DE-
10 FINED.—Section 505(o)(2) of the Federal Food, Drug,
11 and Cosmetic Act (21 U.S.C. 355(o)(2)) is amended by
12 adding at the end the following new subparagraph:

13 “(D) NEW EFFECTIVENESS INFORMA-
14 TION.—The term ‘new effectiveness informa-
15 tion’, with respect to a drug that is or contains
16 a controlled substance for which a listing in any
17 schedule is in effect (on a temporary or perma-
18 nent basis) under section 201 of the Controlled
19 Substances Act, means new information about
20 the effectiveness of the drug, including a new
21 analysis of existing information, derived from—

22 “(i) a clinical trial; an adverse event
23 report; a postapproval study or clinical
24 trial (including a study or clinical trial
25 under paragraph (3));

1 “(ii) peer-reviewed biomedical lit-
 2 erature;

3 “(iii) data derived from the postmar-
 4 ket risk identification and analysis system
 5 under subsection (k); or

6 “(iv) other scientific data determined
 7 to be appropriate by the Secretary.”.

8 (e) CONFORMING AMENDMENTS WITH RESPECT TO
 9 LABELING CHANGES.—Section 505(o)(4) of the Federal
 10 Food, Drug, and Cosmetic Act (21 U.S.C. 355(o)(4)) is
 11 amended—

12 (1) in subparagraph (A)—

13 (A) in the heading, by inserting “OR NEW
 14 EFFECTIVENESS” after “SAFETY”;

15 (B) by striking “safety information” and
 16 inserting “new safety information or new effec-
 17 tiveness information such”; and

18 (C) by striking “believes should be” and
 19 inserting “believes changes should be made to”;

20 (2) in subparagraph (B)(i)—

21 (A) by striking “new safety information”
 22 and by inserting “new safety information or
 23 new effectiveness information”; and

24 (B) by inserting “indications,” after
 25 “boxed warnings,”;

1 (3) in subparagraph (C), by inserting “or new
2 effectiveness information” after “safety informa-
3 tion”; and

4 (4) in subparagraph (E), by inserting “or new
5 effectiveness information” after “safety informa-
6 tion”.

7 (f) RULE OF CONSTRUCTION.—Nothing in the
8 amendments made by this section shall be construed to
9 alter, in any manner, the meaning or application of the
10 provisions of paragraph (3) of section 505(o) of the Fed-
11 eral Food, Drug, and Cosmetic Act (21 U.S.C. 355(o))
12 with respect to the authority of the Secretary of Health
13 and Human Services to require a postapproval study or
14 clinical trial for a purpose specified in clauses (i) through
15 (iii) of subparagraph (B) of such paragraph (3) or para-
16 graph (4) of such section 505(o) with respect to the Sec-
17 retary’s authority to require safety labeling changes.

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