

119TH CONGRESS
1ST SESSION

H. R. 685

To amend the Federal Food, Drug, and Cosmetic Act to prohibit the approval of new abortion drugs, to prohibit investigational use exemptions for abortion drugs, and to impose additional regulatory requirements with respect to previously approved abortion drugs, and for other purposes.

IN THE HOUSE OF REPRESENTATIVES

JANUARY 23, 2025

Mr. LATTA (for himself, Mr. ROUZER, Mr. BRECHEEN, Mr. STRONG, Mrs. MILLER of Illinois, Mr. WEBSTER of Florida, Mr. FINSTAD, Mr. ADERHOLT, Mr. FEENSTRA, Mr. SMITH of New Jersey, Mr. FULCHER, Mr. FLOOD, Mr. MANN, Mr. HARRIS of Maryland, Mr. FONG, Mr. ELLZEY, Mr. WEBER of Texas, Mr. MCCORMICK, Mr. MOOLENAAR, Mr. OGLES, Mr. GUEST, Mr. HIGGINS of Louisiana, Mr. PALMER, Mr. MOORE of North Carolina, Mr. SHREVE, and Mr. LAHOOD) 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 to prohibit the approval of new abortion drugs, to prohibit investigational use exemptions for abortion drugs, and to impose additional regulatory requirements with respect to previously approved abortion drugs, 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,*

1 **SECTION 1. SHORT TITLE.**

2 This Act may be cited as the “Support And Value
3 Expectant Moms and Babies Act of 2025” or the “SAVE
4 Moms and Babies Act of 2025”.

5 **SEC. 2. ABORTION DRUGS PROHIBITED.**

6 (a) IN GENERAL.—Section 505 of the Federal Food,
7 Drug, and Cosmetic Act (21 U.S.C. 355) (as amended by
8 Public Law 117–328) is amended by adding at the end
9 the following:

10 “(aa) ABORTION DRUGS.—

11 “(1) PROHIBITIONS.—The Secretary shall not
12 approve—

13 “(A) any application submitted under sub-
14 section (b) or (j) for marketing an abortion
15 drug; or

16 “(B) grant an investigational use exemp-
17 tion under subsection (i) for—

18 “(i) an abortion drug; or

19 “(ii) any investigation in which the
20 unborn child of a woman known to be
21 pregnant is knowingly destroyed.

22 “(2) PREVIOUSLY APPROVED ABORTION
23 DRUGS.—If an approval described in paragraph (1)
24 is in effect for an abortion drug as of the date of
25 enactment of the Support And Value Expectant
26 Moms and Babies Act of 2025, the Secretary shall—

1 “(A) not approve any labeling change—

2 “(i) to approve the use of such abor-
3 tion drug after 70 days gestation; or

4 “(ii) to approve the dispensing of such
5 abortion drug by any means other than in-
6 person administration by the prescribing
7 health care practitioner;

8 “(B) treat such abortion drug as subject to
9 section 503(b)(1); and

10 “(C) require such abortion drug to be sub-
11 ject to a risk evaluation and mitigation strategy
12 under section 505–1 that at a minimum—

13 “(i) requires health care practitioners
14 who prescribe such abortion drug—

15 “(I) to be certified in accordance
16 with the strategy; and

17 “(II) to not be acting in their ca-
18 pacity as a pharmacist;

19 “(ii) as part of the certification proc-
20 ess referred to in clause (i), requires such
21 practitioners—

22 “(I) to have the ability to assess
23 the duration of pregnancy accurately;

24 “(II) to have the ability to diag-
25 nose ectopic pregnancies;

1 “(III) to have the ability to pro-
2 vide surgical intervention in cases of
3 incomplete abortion or severe bleed-
4 ing;

5 “(IV) to have the ability to en-
6 sure patient access to medical facili-
7 ties equipped to provide blood trans-
8 fusions and resuscitation, if necessary;
9 and

10 “(V) to report any deaths or
11 other adverse events associated with
12 the use of such abortion drug to the
13 Food and Drug Administration and to
14 the manufacturer of such abortion
15 drug, identifying the patient by a non-
16 identifiable reference and the serial
17 number from each package of such
18 abortion drug;

19 “(iii) limits the dispensing of such
20 abortion drug to patients—

21 “(I) in a clinic, medical office, or
22 hospital by means of in-person admin-
23 istration by the prescribing health
24 care practitioner; and

1 “(II) not in pharmacies or any
2 setting other than the health care set-
3 tings described in subclause (I);

4 “(iv) requires the prescribing health
5 care practitioner to give to the patient doc-
6 umentation on any risk of serious com-
7 plications associated with use of such abor-
8 tion drug and receive acknowledgment of
9 such receipt from the patient;

10 “(v) requires all known adverse events
11 associated with such abortion drug to be
12 reported, excluding any individually identi-
13 fiable patient information, to the Food and
14 Drug Administration by the—

15 “(I) manufacturers of such abor-
16 tion drug; and

17 “(II) prescribers of such abortion
18 drug; and

19 “(vi) requires reporting of administra-
20 tion of the abortion drug as required by
21 State law, or in the absence of a State law
22 regarding such reporting, in the same
23 manner as a surgical abortion.

24 “(3) REPORTING ON ADVERSE EVENTS BY
25 OTHER HEALTH CARE PRACTITIONERS.—The Sec-

1 retary shall require all other health care practi-
2 tioners to report to the Food and Drug Administra-
3 tion any adverse events experienced by their patients
4 that are connected to use of an abortion drug, ex-
5 cluding any individually identifiable patient informa-
6 tion.

7 “(4) RULE OF CONSTRUCTION.—Nothing in
8 this section shall be construed to restrict the author-
9 ity of the Federal Government, or of a State, to es-
10 tablish, implement, and enforce requirements and re-
11 strictions with respect to abortion drugs under provi-
12 sions of law other than this section that are in addi-
13 tion to the requirements and restrictions under this
14 section.

15 “(5) DEFINITIONS.—In this section:

16 “(A) The term ‘abortion drug’ means any
17 drug, substance, or combination of drugs or
18 substances that is intended for use or that is in
19 fact used (irrespective of how the product is la-
20 beled) to intentionally kill the unborn child of
21 a woman known to be pregnant, or to inten-
22 tionally terminate the pregnancy of a woman
23 known to be pregnant, with an intention other
24 than—

25 “(i) to produce a live birth;

1 “(ii) to remove a dead unborn child;

2 or

3 “(iii) to treat an ectopic pregnancy.

4 “(B) The term ‘adverse event’ includes
5 each of the following:

6 “(i) A fatality.

7 “(ii) An ectopic pregnancy.

8 “(iii) A hospitalization.

9 “(iv) A blood loss requiring a trans-
10 fusion.

11 “(v) An infection, including endo-
12 metritis, pelvic inflammatory disease, and
13 pelvic infections with sepsis.

14 “(vi) A severe infection.

15 “(C) The term ‘gestation’ means the pe-
16 riod of days beginning on the first day of the
17 last menstrual period.

18 “(D) The term ‘health care practitioner’
19 means any individual who is licensed, reg-
20 istered, or otherwise permitted, by the United
21 States or the jurisdiction in which the indi-
22 vidual practices, to prescribe drugs subject to
23 section 503(b)(1).

24 “(E) The term ‘unborn child’ means an in-
25 dividual organism of the species homo sapiens,

1 beginning at fertilization, until the point of
2 being born alive as defined in section 8(b) of
3 title 1, United States Code.”.

4 (b) ONGOING INVESTIGATIONAL USE.—In the case of
5 any investigational use of a drug pursuant to an investiga-
6 tional use exemption under section 505(i) of the Federal
7 Food, Drug, and Cosmetic Act (21 U.S.C. 355(i)) that
8 was granted before the date of enactment of this Act, such
9 exemption is deemed to be rescinded as of the day that
10 is 3 years after the date of enactment of this Act if the
11 Secretary would be prohibited by section 505(aa)(1)(B) of
12 the Federal Food, Drug, and Cosmetic Act, as added by
13 subsection (a), from granting such exemption as of such
14 day.

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