To provide that the approved application under the Federal Food, Drug, and Cosmetic Act for the drug commonly known as RU-486 is deemed to have been withdrawn, to provide for the review by the Comptroller General of the United States of the process by which the Food and Drug Administration approved such drug, and for other purposes.

IN THE HOUSE OF REPRESENTATIVES

Mr. DEMINT introduced the following bill; which was referred to the Committee on _____

A BILL

- To provide that the approved application under the Federal Food, Drug, and Cosmetic Act for the drug commonly known as RU-486 is deemed to have been withdrawn, to provide for the review by the Comptroller General of the United States of the process by which the Food and Drug Administration approved such drug, and for other purposes.
 - 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 "[RU-486 Approval
5 Suspension and Review Act of 2003]".



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1	SEC. 2. SUSPENSION OF APPROVAL OF DRUG COMMONLY
2	KNOWN AS RU-486; REVIEW AND REPORT BY
3	GENERAL ACCOUNTING OFFICE.
4	(a) IN GENERAL.—Effective upon the expiration of
5	14 days after the date of the enactment of this Act:
6	(1) The approved application under section
7	505(b) of the Federal Food, Drug, and Cosmetic Act
8	for the drug mifepristone (marketed as Mifeprex,
9	and commonly known as RU-486) is deemed to have
10	been withdrawn under section 505(e) of such Act.
11	(2) Such drug shall be considered adulterated
12	for purposes of section 301 of such Act.
13	(b) Review and Report by General Accounting
14	Office.—
15	(1) IN GENERAL.—The Comptroller General of
16	the United States shall review the process by which
17	the Food and Drug Administration approved
18	mifepristone under section 505 of the Federal Food,
19	Drug, and Cosmetic Act and shall determine wheth-
20	er such approval was provided in accordance with
21	such section. The Secretary of Health and Human
22	Services shall ensure that the Comptroller General
23	has full access to all information possessed by the
24	Department of Human Services that relates to such
25	process.



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(2) REPORT.—Not later than 180 days after
 the date of the enactment of this Act, the Comp troller General shall complete the review under para graph (1) and submit to the Congress and the Sec retary of Health and Human Services a report that
 provides the findings of the review.

7 (c) Contingent Reinstatement of Approval of 8 DRUG.—If the report under subsection (b) includes a de-9 termination by the Comptroller General that the approval 10 by the Food and Drug Administration of mifepristone was provided in accordance with section 505 of the Federal 11 12 Food, Drug, and Cosmetic Act, the Secretary of Health 13 and Human Services shall publish such statement in the Federal Register. Effective upon the expiration of 30 days 14 15 after such publication, subsection (a) ceases to have any 16 legal effect.



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