

109TH CONGRESS
2^D SESSION

H. R. 5922

To amend the Federal Food, Drug, and Cosmetic Act to establish additional authorities to ensure the safe and effective use of drugs, to establish whistleblower protections for certain individuals, and for other purposes.

IN THE HOUSE OF REPRESENTATIVES

JULY 27, 2006

Mr. MARKEY 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 establish additional authorities to ensure the safe and effective use of drugs, to establish whistleblower protections for certain individuals, 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 “Swift Approval, Full
5 Evaluation Drug Act” or the “SAFE Drug Act”.

1 **SEC. 2. POSTMARKET STUDIES REGARDING DRUG SAFETY;**
2 **POSTMARKET LABELING CHANGES.**

3 (a) POSTMARKET ORDERS REGARDING STUDIES.—
4 Chapter V of the Federal Food, Drug, and Cosmetic Act
5 (21 U.S.C. 351 et seq.) is amended by inserting after sec-
6 tion 505B the following section:

7 **“SEC. 505C. POSTMARKET STUDIES REGARDING DRUG**
8 **SAFETY.**

9 “(a) POSTMARKET ORDERS.—Within 30 days after
10 receiving evidence of a significant issue regarding the safe-
11 ty or lack of effectiveness of an approved drug, including
12 reports of adverse events, studies conducted or reports re-
13 leased by the Food and Drug Administration, the National
14 Institutes of Health, the Agency for Healthcare Research
15 and Quality or another relevant agency, actions or reports
16 by regulatory agencies in foreign countries, or studies or
17 case reports published in scientific or academic journals,
18 the Secretary, after providing public notice of the signifi-
19 cant safety or effectiveness issue, may order the holder
20 of the approved application to conduct a study or studies
21 to address the issues involved. Each such notice and order
22 shall be published in the Federal Register.

23 “(b) RESTRICTIONS ON USE.—

24 “(1) IN GENERAL.—An order under subsection
25 (a) with respect to an approved drug may, during
26 the period in which the study involved is conducted,

1 establish restrictions on the distribution or use of
2 the drug if the Secretary determines that such re-
3 strictions are necessary to ensure the safe use of the
4 drug during such period.

5 “(2) CERTAIN AUTHORITIES.—Restrictions that
6 may be established by the Secretary under para-
7 graph (1) with respect to a drug include the fol-
8 lowing:

9 “(A) Restricting distribution to certain fa-
10 cilities or physicians with special training or ex-
11 perience.

12 “(B) Conditioning distribution on the per-
13 formance of specified medical procedures.

14 “(C) Restricting direct-to-consumer adver-
15 tisements for the drug.

16 “(3) TERMINATION.—The Secretary may, on
17 the basis of the results of the study or other evi-
18 dence, continue the restrictions under paragraph (1),
19 terminate restrictions established or establish dif-
20 ferent restrictions, as necessary to ensure the safe
21 use of the drug. The Secretary shall notify the spon-
22 sor of the decision to extend, terminate or change
23 the restrictions no later than 30 days after the date
24 on which the results of the study involved are sub-
25 mitted to the Secretary.

1 “(c) DEFINITION.—For purposes of this section, the
2 term ‘approved drug’ means a drug for which an approved
3 application under section 505 is in effect or for which a
4 biologics license under section 351 of the Public Health
5 Service Act is in effect.”.

6 (b) NEW DRUG APPLICATIONS; POSTMARKET STUD-
7 IES PURSUANT TO ACCELERATED APPROVAL.—Section
8 505 of the Federal Food, Drug, and Cosmetic Act (21
9 U.S.C. 355) is amended by adding at the end the following
10 subsection:

11 “(o)(1) The Secretary shall amend subpart H of part
12 314 of title 21, Code of Federal Regulations, to establish
13 the following policies:

14 “(A) As a condition of the approval under such
15 subpart of a new drug on or after the date of the
16 enactment of the SAFE Drug Act, the Secretary
17 shall require that one or more postmarket studies of
18 the drug be conducted.

19 “(B) The Secretary may not approve the appli-
20 cation involved unless—

21 “(i) the sponsor has submitted to the Sec-
22 retary the protocols for each such study;

23 “(ii) the Secretary has approved the proto-
24 cols; and

1 “(iii) the Secretary and the sponsor have
2 agreed on a timeframe, including designated
3 milestones, for the prompt completion of the
4 study, which timeframe assumes due diligence
5 by the sponsor in conducting the study.

6 “(C) The Secretary shall require that, after the
7 application is approved under such subpart and the
8 drug enters commercial distribution, the drug be
9 marketed in accordance with the following:

10 “(i) The distribution and use of the drug
11 shall be restricted in accordance with such sub-
12 part.

13 “(ii) Until the Secretary determines that
14 the sponsor has fulfilled its commitments under
15 subparagraphs (A) and (B), the labeling of the
16 drug shall bear—

17 “(I) a statement that the Food and
18 Administration is requiring a study or
19 studies to confirm the safety and effective-
20 ness of the drug; and

21 “(II) a statement providing a clear
22 and concise summary of the outstanding
23 issues or questions to be addressed in such
24 required studies.

1 “(iii) Until the Secretary determines that
2 the sponsor has fulfilled its commitments under
3 subparagraphs (A) and (B), the label of the
4 drug shall bear a statement providing as fol-
5 lows: ‘This product received conditional ap-
6 proval from the FDA under its accelerated ap-
7 proval process. It will not receive full approval
8 until completion of further testing to confirm
9 safety and/or efficacy. For further information
10 please contact your physician.’.

11 “(iv) Direct-to-consumer advertisements
12 for the drug shall be restricted until—

13 “(I) the Secretary determines that the
14 sponsor has fulfilled its commitments
15 under subparagraphs (A) and (B); and

16 “(II) the drug has been approved
17 under such part 314 independently of sub-
18 part H of such part.

19 “(2) The Secretary shall amend part 314 of title 21,
20 Code of Federal Regulations, to establish the following
21 policies regarding postmarket studies that, on or after the
22 date of the enactment of the SAFE Drug Act, are required
23 pursuant to approval of a new drug under subpart H of
24 such part or under section 506(b)(2) of this Act:

1 “(A) If a required study is not completed by
2 two years after the date on which the new drug was
3 so approved, the Secretary shall, promptly after the
4 expiration of such period, convene a public meeting
5 of the appropriate advisory committee to review the
6 progress of the study. Such review shall include the
7 assessment of compliance with the timeframe for
8 prompt completion of the study as agreed to by the
9 company and the Secretary under paragraph
10 (1)(B)(iii) or under section 506(b)(2) (as the case
11 may be), the quality of study conduct, rates of en-
12 rollment overall and by institution, the barriers to
13 progress, and whether the sponsor is acting with due
14 diligence. At the meeting, the advisory committee
15 shall determine whether it is in the best interest of
16 the public to allow the sponsor to continue mar-
17 keting the drug until the study is completed or
18 whether it is in the best interest of the public to sus-
19 pend the commercial marketing of the drug until the
20 study is completed.

21 “(B) If the drug was approved on the basis of
22 animal efficacy data because human efficacy studies
23 are not ethical or feasible, the holder of the ap-
24 proved application shall conduct studies when ethical

1 and feasible to verify and describe clinical benefit
2 and to assess the product’s safety.

3 “(C) If the results of a completed study that
4 was so required are inconclusive or the risk-to-ben-
5 efit profile cannot be positively established, the Sec-
6 retary shall withdraw the product for commercial
7 distribution. If a required study is not completed by
8 eight years after the date of approval, the results of
9 the study are presumed to be inconclusive. The prod-
10 uct may only be made available to patients who—

11 “(i) according to a health care professional
12 have previously benefitted from the product;
13 and

14 “(ii) have signed a statement of informed
15 consent stating that they have received notice
16 from the sponsor that the results of completed
17 studies on the product have proven to be incon-
18 clusive and/or the risk to benefit profile can not
19 be positively established. The document shall in-
20 clude the risks of continuing the product.

21 Otherwise the product shall be unavailable except in
22 a research setting until the product can meet the
23 same standard of safety and effectiveness that exists
24 for full approval.”.

1 (c) ENFORCEMENT REGARDING POSTMARKET STUD-
2 IES.—

3 (1) MISBRANDING.—Section 502 of the Federal
4 Food, Drug, and Cosmetic Act (21 U.S.C. 352) is
5 amended by adding at the end the following:

6 “(x) If it is a drug with respect to which there is
7 a failure to comply with a requirement under section
8 505(o)(1), an order under section 505C, or a requirement
9 under section 506(b)(2).”.

10 (2) CIVIL PENALTIES.—Section 303 of the Fed-
11 eral Food, Drug, and Cosmetic Act (21 U.S.C. 331)
12 is amended by adding at the end the following:

13 “(g)(1)(A) Any person who violates section 301(a),
14 301(b), or 301(c) by reason of section 502(x) shall be sub-
15 ject to a civil penalty of not more than 100 percent of
16 the gross profits received by the sponsor from sales of the
17 drug, or \$1,000,000, whichever is greater, subject to sub-
18 paragraph (B).

19 “(B) If any harm to a consumer occurs as a result
20 of a violation referred to in subparagraph (A), the person
21 involved may be subject to a civil penalty of not more than
22 an amount equal to 300 percent of the gross profits re-
23 ceived by the person, or \$3,000,000, whichever is greater.

24 “(2) Any person who fails to act with due diligence
25 to complete a postmarket study required under section

1 506(b)(2), or under subpart H of part 314 of title 21,
2 Code of Federal Regulations, shall be subject to a civil
3 penalty of not more than the amount that applies under
4 subparagraph (A) or subparagraph (B) of paragraph (1)
5 (as the case may be) for a violation referred to in such
6 paragraph. The Secretary shall by regulation define the
7 term ‘due diligence’ for purposes of this paragraph.

8 “(3) The provisions of paragraphs (3) through (5)
9 of subsection (f) apply to a civil penalty under subpara-
10 graph (A) or (B) of paragraph (1) or under paragraph
11 (2) to the same extent and in the same manner as such
12 provisions apply to a civil penalty under paragraph (1) or
13 (2) of such subsection.”.

14 (d) POSTMARKET LABELING CHANGES.—Section
15 502 of the Federal Food, Drug, and Cosmetic Act, as
16 amended by subsection (c)(1) of this section, is amended
17 by adding at the end the following:

18 “(y) If it is a drug and the manufacturer of the
19 drug or product fails to make changes to a product’s
20 labeling in compliance with an order of the Sec-
21 retary, issued on the basis of clinical evidence (in-
22 cluding studies submitted to the Secretary, an anal-
23 ysis of adverse events reports, studies conducted or
24 reports released by the Food and Drug Administra-
25 tion, the National Institutes of Health, the Agency

1 for Healthcare Research and Quality or another rel-
2 evant agency, actions or reports by regulatory agen-
3 cies in foreign countries, or studies published in sci-
4 entific or academic journals), that the labeling of the
5 drug be modified to include specific wording re-
6 quired by the Secretary to ensure the safe and effec-
7 tive use of the drug.”.

8 (e) **RULE OF CONSTRUCTION REGARDING CERTAIN**
9 **PEDIATRIC STUDIES.**—The amendments made by this
10 section establish authorities in addition to, and not in lieu
11 of—

12 (1) the program under section 505A of the
13 Federal Food, Drug, and Cosmetic Act; and

14 (2) authorities under section 505B of such Act.

15 **SEC. 3. WHISTLEBLOWER PROTECTIONS.**

16 (a) **PROHIBITION.**—It shall be unlawful for any per-
17 son to discharge, demote, suspend, reprimand, investigate,
18 or take or fail to take any other personnel action that in
19 any manner discriminates against any covered individual,
20 or in any other manner discriminate against any covered
21 individual (including by a denial, suspension, or revocation
22 of a security clearance or by any other security access de-
23 termination, or by denial of award of a Federal contract
24 or subcontract), or to threaten or recommend the dis-
25 charge, demotion, suspension, reprimand, investigation,

1 other personnel action (or rejection of such action) that
2 in any manner discriminates against any covered indi-
3 vidual, or other manner of discrimination if such action,
4 discrimination, or recommendation is due, in whole or in
5 part, to any lawful act done, perceived to have been done,
6 or intended to be done by the covered individual—

7 (1) to provide information, cause information to
8 be provided, or otherwise assist in an investigation
9 or proceeding regarding any conduct which the cov-
10 ered individual reasonably believes constitutes evi-
11 dence of a violation of any law, rule, or regulation,
12 a substantial and specific threat to public health or
13 safety, abuse of authority, or fraud, waste, or mis-
14 management of public funds, if the information or
15 assistance is provided to or the investigation or pro-
16 ceeding is conducted by—

17 (A) a Federal, State, or local regulatory or
18 law enforcement agency (including an office of
19 Inspector General under the Inspector General
20 Act of 1978);

21 (B) any Member of Congress, any com-
22 mittee of Congress, or the Government Ac-
23 countability Office;

24 (C) any person with supervisory or mana-
25 gerial authority over the covered individual (or

1 any other person who has the authority to in-
2 vestigate, discover, or terminate misconduct); or

3 (D) a potential witness to or other person
4 affected by or aware of the conduct described in
5 this section who has the authority to inves-
6 tigate, discover, or terminate misconduct;

7 (2) to file, cause to be filed, testify, participate
8 in, or otherwise assist in a proceeding or action filed
9 or about to be filed relating to an alleged violation
10 of any law, rule, or regulation; or

11 (3) to refuse to violate or assist in the violation
12 of any law, rule, or regulation.

13 (b) ENFORCEMENT ACTION.—

14 (1) IN GENERAL.—A covered individual who al-
15 leges discharge or other discrimination by any per-
16 son in violation of subsection (a) may seek relief
17 under subsection (c) by—

18 (A) filing a complaint with the Secretary of
19 Labor; or

20 (B) if the Secretary has not issued a final
21 decision within 180 days after the filing of the
22 complaint and there is no showing that such
23 delay is due to the bad faith of the claimant,
24 bringing an action at law or equity for de novo
25 review in the appropriate district court of the

1 United States, which shall have jurisdiction
2 over such an action without regard to the
3 amount in controversy.

4 (2) PROCEDURE.—

5 (A) IN GENERAL.—An action under para-
6 graph (1)(A) shall be governed under the rules
7 and procedures set forth in section 42121(b) of
8 title 49, United States Code.

9 (B) EXCEPTION.—Notification made under
10 section 42121(b)(1) of title 49, United States
11 Code, shall be made—

12 (i) to the person named in the com-
13 plaint; and

14 (ii) to the person's employer.

15 (C) BURDENS OF PROOF.—An action
16 brought under paragraph (1)(B) shall be gov-
17 erned by the legal burdens of proof set forth in
18 section 42121(b) of title 49, United States
19 Code.

20 (D) STATUTE OF LIMITATIONS.—An action
21 under paragraph (1) shall be commenced not
22 later than 6 years after the date on which the
23 violation occurs.

24 (c) REMEDIES.—

1 (1) IN GENERAL.—A covered individual pre-
2 vailing in any action under subsection (b) shall be
3 entitled to all relief appropriate to make the covered
4 individual whole.

5 (2) DAMAGES.—Relief for any action under
6 subsection (b) shall include—

7 (A) reinstatement with the same seniority
8 status and employment grade or pay level (or
9 the equivalent) that the covered individual
10 would have had, but for the discrimination;

11 (B) compensatory damages, including the
12 amount of any back pay, with interest;

13 (C) compensation for any special damages
14 sustained as a result of the discrimination, in-
15 cluding litigation costs, expert witness fees, and
16 reasonable attorney fees; and

17 (D) punitive damages in an amount not to
18 exceed the greater of 3 times the amount of any
19 monetary damages awarded under this section
20 (apart from this paragraph) or \$5,000,000.

21 (d) RIGHTS RETAINED BY COVERED INDIVIDUAL.—
22 Nothing in this section shall be deemed to diminish the
23 rights, privileges, or remedies of any covered individual
24 under any Federal or State law, or under any collective
25 bargaining agreement. The rights and remedies in this

1 section may not be waived by any agreement, policy, form,
2 or condition of employment.

3 (e) NOTIFICATION.—The provisions of this section
4 shall be prominently posted in any place of employment
5 to which this section applies.

6 (f) DEFINITIONS.—For purposes of this section:

7 (1) The term “covered individual” means an
8 employee of the Food and Drug Administration.

9 (2) The term “lawful” means not specifically
10 prohibited by law.

11 **SEC. 4. RIGHT TO PUBLISH.**

12 Subchapter E of chapter V of the Federal Food,
13 Drug, and Cosmetic Act (21 U.S.C. 360bbb et seq.) is
14 amended by adding at the end the following section:

15 **“SEC. 565. RIGHT TO PUBLISH.**

16 “Officers and employees of the Food and Drug Ad-
17 ministration, and individuals sponsored by such Adminis-
18 tration, may publish in peer-reviewed journals and other
19 scientific publications, and make oral presentations at pro-
20 fessional society meetings and other meetings of their
21 peers, unless publication or presentation of the data is
22 subject to Federal export control or national security laws
23 or regulations, or is proprietary information. The right to
24 publish or present such data cannot be waived by any
25 agreement, policy, form or condition of employment.”.

1 **SEC. 5. BIENNIAL REPORTS ON APPROVED APPLICATIONS**
2 **SUPPORTED BY NON-INFERIORITY STUDIES.**

3 Section 505 of the Federal Food, Drug, and Cosmetic
4 Act, as amended by section 2(b) of this Act, is amended
5 by adding at the end the following subsection:

6 “(p) BIENNIAL REPORTS ON APPROVED APPLICA-
7 TIONS SUPPORTED BY NON-INFERIORITY STUDIES.—The
8 Secretary shall submit to the Congress biennial reports on
9 approved applications under subsection (b) (including sup-
10 plemental applications) that have been supported by data
11 from one or more non-inferiurity studies. For each such
12 application, the report shall include the following informa-
13 tion:

14 “(1) The name of the drug listed in application.

15 “(2) The name of the sponsor.

16 “(3) The date of the approval.

17 “(4) The name of each drug used as an active
18 control in the non-inferiurity studies used to support
19 the application.

20 “(5) The indication studied.

21 “(6) The primary and secondary endpoints of
22 the non-inferiurity studies.

23 “(7) The margins used in such studies.

24 “(8) The explanation required by section
25 314.126(b)(2)(iv) of title 21, Code of Federal Regu-

1 lations, as to why the drugs should be considered ef-
2 fective in the study.”.

3 **SEC. 6. BIENNIAL REPORTS REGARDING POSTMARKET**
4 **STUDIES.**

5 Section 505 of the Federal Food, Drug, and Cosmetic
6 Act, as amended by section 5 of this Act, is amended by
7 adding at the end the following subsection:

8 “(q) BIENNIAL REPORTS REGARDING POSTMARKET
9 STUDIES.—The Secretary shall submit to the Congress bi-
10 annual reports that provide the following information:

11 “(1) The number of enforcement actions taken
12 to ensure that sponsors are complying with the re-
13 quirements to complete postmarketing studies under
14 subsection (o) and sections 505C and 506(b)(2), to-
15 gether with a description of each such action.

16 “(2) The measures taken by the Secretary to
17 establish a system to ensure effective monitoring of
18 the status of all such postmarketing studies, to-
19 gether with a description of the status of that sys-
20 tem.

21 “(3) The measures taken by the Secretary to
22 develop a system to track information about ongoing
23 postmarketing safety issues, together with a descrip-
24 tion of the status of that system.”.

○