# <sup>108TH CONGRESS</sup> IST SESSION H.R. 2917

To amend the Federal Food, Drug, and Cosmetic Act with respect to the safety of genetically engineered foods, and for other purposes.

#### IN THE HOUSE OF REPRESENTATIVES

#### JULY 25, 2003

Mr. KUCINICH (for himself, Mr. DEFAZIO, Mr. SANDERS, Ms. LEE, Mr. CON-YERS, Mr. OLVER, Mr. GEORGE MILLER of California, Mr. HONDA, Mr. ACEVEDO-VILÁ, Mr. GUTIERREZ, Mr. NADLER, Mr. OWENS, Ms. VELÁZQUEZ, Ms. WATERS, Ms. WATSON, and Ms. WOOLSEY) 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 the safety of genetically engineered foods, 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 "Genetically Engineered
- 5 Food Safety Act".

#### 6 SEC. 2. FINDINGS.

7 The Congress finds as follows:

(1) Genetic engineering is an artificial gene
 transfer process wholly different from traditional
 breeding.

4 (2) Genetic engineering can be used to produce
5 new versions of virtually all plant and animal foods.
6 Thus, within a short time, the food supply could
7 consist almost entirely of genetically engineered
8 products.

9 (3) This conversion from a food supply based 10 on traditionally bred organisms to one based on or-11 ganisms produced through genetic engineering could 12 be one of the most important changes in our food 13 supply in this century.

14 (4) Genetically engineered foods present new
15 issues of safety that have not been adequately stud16 ied.

17 (5) The Congress has previously required that
18 food additives be analyzed for their safety prior to
19 their placement on the market.

20 (6) Adding new genes into a food should be
21 considered adding a food additive, thus requiring an
22 analysis of safety factors.

(7) Federal agencies have failed to uphold congressional intent of the Food Additives Amendment
of 1958 by allowing genetically engineered foods to

be marketed, sold and otherwise used without re quiring pre-market safety testing addressing their
 unique characteristics.

4 (8) The food additive process gives the Food
5 and Drug Administration discretion in applying the
6 safety factors that are generally recognized as ap7 propriate to evaluate the safety of food and food in8 gredients.

9 (9) Given the consensus among the scientific 10 community that genetic engineering can potentially 11 introduce hazards, such as allergens or toxins, ge-12 netically engineered foods need to be evaluated on a 13 case-by-case basis and cannot be presumed to be 14 generally recognized as safe.

15 SEC. 3. FEDERAL DETERMINATION OF SAFETY OF GENETI-

16 CALLY ENGINEERED FOOD; REGULATION AS17 FOOD ADDITIVE.

18 (a) INCLUSION IN DEFINITION OF FOOD ADDI19 TIVE.—Section 201 of the Federal Food, Drug, and Cos20 metic Act (21 U.S.C. 321) is amended—

21 (1) in paragraph (s), by adding after and below
22 subparagraph (6) the following sentence:

23 "Such term includes the different genetic constructs, pro-24 teins of such constructs, vectors, promoters, marker sys-25 tems, and other appropriate terms that are used or cre-

ated as a result of the creation of a genetically engineered 1 2 food (as defined in paragraph (nn)), other than a genetic 3 construct, protein, vector, promoter, or marker system or 4 other appropriate term for which an application under sec-5 tion 505 or 512 has been filed. For purposes of this Act, the term 'genetic food additive' means a genetic construct, 6 7 protein, vector, promoter, or marker system or other ap-8 propriate term that is so included."; and

10 "(nn)(1) The term 'genetically engineered food'
11 means food that contains or was produced with a geneti12 cally engineered material.

(2) by adding at the end the following:

13 "(2) The term 'genetically engineered material' 14 means material derived from any part of a genetically en-15 gineered organism, without regard to whether the altered 16 molecular or cellular characteristics of the organism are 17 detectable in the material.

18 "(3) The term 'genetically engineered organism'19 means—

"(A) an organism that has been altered at the
molecular or cellular level by means that are not
possible under natural conditions or processes (including but not limited to recombinant DNA and
RNA techniques, cell fusion, microencapsulation,
macroencapsulation, gene deletion and doubling, in-

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troducing a foreign gene, and changing the positions
 of genes), other than a means consisting exclusively
 of breeding, conjugation, fermentation, hybridiza tion, in vitro fertilization, tissue culture, or
 mutagenesis; and

6 "(B) an organism made through sexual or asex7 ual reproduction (or both) involving an organism de8 scribed in clause (A), if possessing any of the altered
9 molecular or cellular characteristics of the organism
10 so described.

11 "(4) For purposes of subparagraph (1), a food shall 12 be considered to have been produced with a genetically engineered material if the organism from which the food is 13 14 derived has been injected or otherwise treated with a ge-15 netically engineered material (except that the use of manure as a fertilizer for raw agricultural commodities may 16 17 not be construed to mean that such commodities are pro-18 duced with a genetically engineered material).".

19 (b) Petition To Establish Safety.—

(1) DATA IN PETITION.—Section 409(b)(2)(E)
of the Federal Food, Drug, and Cosmetic Act (21
U.S.C. 348(b)(2)(E)) is amended by adding at the
end the following sentence: "In the case of a genetic
food additive, such reports shall include all data that
was collected or developed pursuant to the investiga-

1	tions, including data that does not support the claim
2	of safety for use.".
3	(2) Notices; public availability of infor-
4	MATION.—Section 409(b)(5) of the Federal Food,
5	Drug, and Cosmetic Act (21 U.S.C. 348(b)(5)) is
6	amended—
7	(A) by striking "(5)" and inserting
8	"(5)(A)"; and
9	(B) by adding at the end the following sub-
10	paragraphs:
11	"(B) In the case of a genetic food additive:
12	"(i) Promptly after providing the notice under
13	subparagraph (A), the Secretary shall make avail-
14	able to the public all reports and data described in
15	paragraph $(2)(E)$ that are contained in the petition
16	involved, and all other information in the petition to
17	the extent that the information is relevant to a de-
18	termination of the safety for use of the additive.
19	"(ii) Such notice shall state whether any infor-
20	mation in the petition is not being made available to
21	the public because the Secretary has made a deter-
22	mination that the information does not relate to the
23	safety for use of the additive. Any person may peti-
24	tion the Secretary for a reconsideration of such a de-
25	termination.

1 "(C) In the case of genetic food additives:

2 "(i) The Secretary shall maintain and make 3 available to the public through telecommunications a 4 list of petitions that are pending under this sub-5 section and a list of petitions for which regulations 6 under subsection (c)(1)(A) have been established. 7 Such list shall include information on the additives 8 involved, including the source of the additives, and 9 including any information received by the Secretary 10 pursuant to clause (ii).

11 "(ii) If a regulation is in effect under sub-12 section (c)(1)(A) for a genetic food additive, any 13 person who manufactures such additive for commer-14 cial use shall submit to the Secretary a notification 15 of any knowledge of data that relate to the adverse 16 health effects of the additive, when knowledge is ac-17 quired by the person after the date on which the 18 regulation took effect. If the manufacturer is in pos-19 session of the data, the notification shall include the 20 data. The Secretary shall by regulation establish the 21 scope of the responsibilities of manufacturers under 22 this clause, including such limits on the responsibil-23 ities as the Secretary determines to be appropriate.".

24 (3) EFFECTIVE DATE OF REGULATION REGARD25 ING SAFE USE; OPPORTUNITY FOR PUBLIC COM-

MENT.—Section 409(c)(2) of the Federal Food,
 Drug, and Cosmetic Act (21 U.S.C. 348(c)(2)) is
 amended—

4 (A) by striking "(2)" and inserting 5 "(2)(A)"; and

6 (B) by adding at the end the following sub-7 paragraph:

8 "(B)(i) In the case of a genetic food additive, an 9 order under paragraph (1)(A) may not be issued regarding 10 the petition involved before the expiration of the applicable period under clause (ii). During such period, and con-11 12 tinuing until an order under paragraph (1) is issued, the 13 Secretary shall provide interested persons an opportunity to submit to the Secretary comments on the petition. In 14 15 publishing such notice, the Secretary shall inform the public of such opportunity. 16

17 "(ii) For purposes of clause (i), the applicable period 18 under this clause regarding a petition is the 30-day period 19 beginning on the date on which the Secretary has under 20subparagraph (B)(i) of subsection (b)(5) made informa-21 tion available to the public regarding the petition, except 22 that, if under subparagraph (B)(ii) of such subsection the 23 Secretary finds in favor of a person who files for reconsid-24 eration (relating to a determination by the Secretary that 25 information does not relate to safety), such 30-day period

is extended by an additional period of 30-days. For pur poses of the preceding sentence, a discrete 30-day exten sion applies to each such reconsideration for which the
 Secretary finds in favor of the person filing for reconsider ation.".

6 (4) CONSIDERATION OF CERTAIN FACTORS.—
7 Section 409(c) of the Federal Food, Drug, and Cos8 metic Act (21 U.S.C. 348(c)) is amended by adding
9 at the end the following paragraph:

"(6) In the case of a genetic food additive, the factors
considered by the Secretary regarding safety for use shall
include (but not be limited to) the results of the following
analyses:

14 "(A) Allergenicity effects resulting from the15 added proteins, including proteins not found in the16 food supply.

"(B) Pleiotropic effects. The Secretary shall require tests to determine the potential for such effects (using molecular characterization, biochemical
characterization, mRNA profiling, or other techniques, or as appropriate, combinations of such techniques).

23 "(C) Appearance of new toxins or increased lev-24 els of existing toxins.

"(D) Changes in the functional characteristics
 of food.

3 "(E) Changes in the levels of important nutri-4 ents.

5 "(F) Changes in the levels of anti-nutrients.".
6 (5) CERTAIN TESTS.—Section 409(c) of the
7 Federal Food, Drug, and Cosmetic Act, as amended
8 by paragraph (4), is amended by adding at the end
9 the following paragraph:

10 "(7) In the case of genetic food additives:

11 "(A) If a genetic food additive is a protein from 12 a commonly or severely allergenic food, the Sec-13 retary may not establish a regulation under para-14 graph (1)(A) if the petition under subsection (b)(1)15 fails to include full reports of investigations that used serum or skin tests (or other advanced tech-16 17 niques) on a sensitive population to determine 18 whether such additive is commonly or severely aller-19 genic.

"(B)(i) If a genetic food additive is a protein
that has not undergone the investigations described
in subparagraph (A), the Secretary may not establish a regulation under paragraph (1)(A) if the petition under subsection (b)(1) fails to include full reports of investigations that used the best available

biochemical and physiological protocols to evaluate
 whether it is likely that the protein involved is an allergen.

4 "(ii) For purposes of clause (i), the Secretary
5 shall by regulation determine the best available bio6 chemical and physiological protocols. In carrying out
7 rulemaking under the preceding sentence, the Sec8 retary shall consult with the Director of the Na9 tional Institutes of Health.".

10 (6) PROHIBITED ADDITIVES.—Section 409(c) of
11 the Federal Food, Drug, and Cosmetic Act, as
12 amended by paragraph (5), is amended by adding at
13 the end the following paragraph:

14 "(8) In the case of a genetic food additive, the Sec15 retary may not establish a regulation under paragraph
16 (1)(A) if—

17 "(A) the additive is a protein and a report of
18 an investigation finds that the additive is likely to be
19 commonly or severely allergenic;

"(B) the additive is a protein and a report of
an investigation that uses a protocol described in
paragraph (7)(B) fails to find with reasonable certainty that the additive is unlikely to be an allergen;
or

"(C) effective June 1, 2006, a selective marker
is used with respect to the additive, the selective
marker will remain in the food involved when the
food is marketed, and the selective marker inhibits
the function of one or more antibiotics.".

6 (7) ADDITIONAL PROVISIONS.—Section 409(c)
7 of the Federal Food, Drug, and Cosmetic Act, as
8 amended by paragraph (6), is amended by adding at
9 the end the following paragraph:

10 ((9)(A) In determining the safety for use of genetic food additives, the Secretary may (directly or through con-11 12 tract) conduct investigations of such additives for purposes of supplementing the information provided to the 13 Secretary pursuant to petitions under subsection (b)(1). 14 15 "(B) To provide the Congress with a periodic independent, external review of the Secretary's formulation of 16 the approval process under paragraph (1)(A) that relates 17 to genetic food additives, the Secretary shall enter into 18 19 an agreement with the Institute of Medicine. Such agree-20 ment shall provide that, if the Institute of Medicine has 21 any concerns regarding the approval process, the Institute 22 of Medicine will submit to the Congress a report describ-

23 ing such concerns.".

(c) REGULATION ISSUED ON SECRETARY'S INITIA TIVE.—Section 409(d) of the Federal Food, Drug, and
 Cosmetic Act (21 U.S.C. 348(d)) is amended—

4 (1) by striking "(d) The Secretary" and insert5 ing "(d)(1) Subject to paragraph (2), the Sec6 retary"; and

7 (2) by adding at the end the following para-8 graph:

9 "(2) The provisions of subsections (b) and (c) that 10 expressly reference genetic food additives apply with re-11 spect to a regulation proposed by the Secretary under 12 paragraph (1) to the same extent and in the same manner 13 as such provisions apply with respect to a petition filed 14 under subsection (b)(1).".

(d) CIVIL PENALTIES.—Section 303 of the Federal
Food, Drug, and Cosmetic Act (21 U.S.C. 333) is amended by adding at the end the following subsection:

18 "(h)(1) With respect to a violation of section 301(a), 19 301(b), or 301(c) involving the adulteration of food by rea-20 son of failure to comply with the provisions of section 409 21 that relate to genetic food additives, any person engaging 22 in such a violation shall be liable to the United States for 23 a civil penalty in an amount not to exceed \$100,000 for 24 each such violation. "(2) Paragraphs (3) through (5) of subsection (g)
 apply with respect to a civil penalty under paragraph (1)
 of this subsection to the same extent and in the same man ner as such paragraphs (3) through (5) apply with respect
 to a civil penalty under paragraph (1) or (2) of subsection
 (g).".

7 (e) CITIZEN SUITS.—Chapter III of the Federal
8 Food, Drug, and Cosmetic Act (21 U.S.C. 331 et seq.)
9 is amended by adding at the end the following section:
10 "CITIZEN SUITS REGARDING GENETIC FOOD ADDITIVES

"SEC. 311. (a) IN GENERAL.—Except as provided in
subsection (c), any person may on his or her behalf commence a civil action in an appropriate district court of the
United States against—

15 "(1) a person who is alleged to have engaged in 16 a violation of section 301(a), 301(b), or 301(c) in-17 volving the adulteration of food by reason of failing 18 to comply with the provisions of section 409 that re-19 late to genetic food additives; or

"(2) the Secretary where there is alleged a failure of the Secretary to perform any act or duty
under section 409 that relates to such additives and
is not discretionary.

24 "(b) RELIEF.—In a civil action under subsection (a),
25 the district court involved may, as the case may be—

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1	((1) enforce the compliance of a person with
2	the applicable provisions referred to paragraph $(1)$
3	of such subsection; or
4	((2)) order the Secretary to perform an act or
5	duty referred to in paragraph (2) of such subsection.
6	"(c) LIMITATIONS.—
7	"(1) NOTICE TO SECRETARY.—A civil action
8	may not be commenced under subsection $(a)(1)$ prior
9	to 60 days after the plaintiff has provided to the
10	Secretary notice of the violation involved.
11	"(2) Relation to actions of secretary.—
12	A civil action may not be commenced under sub-
13	section $(a)(2)$ if the Secretary has commenced and
14	is diligently prosecuting a civil or criminal action in
15	a district court of the United States to enforce com-
16	pliance with the applicable provisions referred to in
17	subsection $(a)(1)$ .
18	"(d) Right of Secretary To Intervene.—In any
19	civil action under subsection (a), the Secretary, if not a
20	party, may intervene as a matter of right.
21	"(e) Award of Costs; Filing of Bond.—In a civil
22	action under subsection (a), the district court involved
23	may award costs of litigation (including reasonable attor-
24	ney and expert witness fees) to any party whenever the
25	court determines such an award is appropriate. The court

may, if a temporary restraining order or preliminary in junction is sought, require the filing of a bond or equiva lent security in accordance with the Federal Rules of Civil
 Procedure.

5 "(f) SAVINGS PROVISION.—This section does not re-6 strict any right that a person (or class of persons) may 7 have under any statute or common law to seek enforce-8 ment of the provisions referred to subsection (a)(1), or to 9 seek any other relief (including relief against the Sec-10 retary).".

11 (f) RULE OF CONSTRUCTION.—With respect to sec-12 tion 409 of the Federal Food, Drug, and Cosmetic Act 13 as amended by this section, compliance with the provisions of such section 409 that relate to genetic food additives 14 15 does not constitute an affirmative defense in any cause of action under Federal or State law for personal injury 16 resulting in whole or in part from a genetic food additive. 17 18 SEC. 4. USER FEES REGARDING DETERMINATION OF SAFE-

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#### TY OF GENETIC FOOD ADDITIVES.

20 Chapter IV of the Federal Food, Drug, and Cosmetic
21 Act (21 U.S.C. 341 et seq.) is amended by inserting after
22 section 409 the following section:

23 "USER FEES REGARDING SAFETY OF GENETIC FOOD

24 ADDITIVES

25 "SEC. 409A. (a) IN GENERAL.—In the case of ge26 netic food additives, the Secretary shall in accordance with
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1	this section assess and collect a fee on each petition that
2	is filed under section $409(b)(1)$ . The fee shall be collected
3	from the person who submits the petition, is due upon sub-
4	mission of the petition, and shall be assessed in an amount
5	determined under subsection (c). This section applies as
6	of the first fiscal year that begins after the date of promul-
7	gation of the final rule required in section 6 of the Geneti-
8	cally Engineered Food Safety Act (referred to in this sec-
9	tion as the 'first applicable fiscal year').
10	"(b) Purpose of Fees.—
11	"(1) IN GENERAL.—The purposes of fees under
12	subsection (a) are as follows:
13	"(A) To defray increases in the costs of
14	the resources allocated for carrying out section
15	409 for the first applicable fiscal year over the
16	costs of carrying out such section for the pre-
17	ceding fiscal year, other than increases that are
18	not attributable to the responsibilities of the
19	Secretary with respect to genetic food additives.
20	"(B) To provide for a program of basic
21	and applied research on the safety of genetic
22	food additives (to be carried out by the Com-
23	missioner). The program shall address funda-
24	mental questions and problems that arise re-
25	peatedly during the process of reviewing peti-

1	tions under section $409(b)(1)$ with respect to
2	genetic food additives, and shall not directly
3	support the development of new genetically en-
4	gineered foods.
5	"(2) Allocations by secretary.—Of the
6	total fee revenues collected under subsection (a) for
7	a fiscal year, the Secretary shall reserve and ex-
8	pend—
9	"(A) 95 percent for the purpose described
10	in paragraph $(1)(A)$ and
11	"(B) 5 percent for the purpose described
12	in paragraph (1)(B).
13	"(3) CERTAIN PROVISIONS REGARDING IN-
14	CREASED ADMINISTRATIVE COSTS.—With respect to
15	fees under subsection (a):
16	"(A) Increases referred to in paragraph
17	(1)(A) include the costs of the Secretary in pro-
18	viding for investigations under section
19	409(c)(9)(A).
20	"(B) Increases referred to in paragraph
21	(1)(A) include increases in costs for an addi-
22	tional number of full-time equivalent positions
23	in the Department of Health and Human Serv-
24	ices to be engaged in carrying out section 409
25	with respect to genetic food additives.

1 "(c) TOTAL FEE REVENUES; INDIVIDUAL FEE 2 AMOUNTS.—The total fee revenues collected under sub-3 section (a) for a fiscal year shall be the amounts appro-4 priated under subsection (f)(2) for such fiscal year. Indi-5 vidual fees shall be assessed by the Secretary on the basis of an estimate by the Secretary of the amount necessary 6 7 to ensure that the sum of the fees collected for such fiscal 8 year equals the amount so appropriated. In assessing the 9 individual fees, the Secretary shall by regulation provide 10 for the assessment of reduced fee amounts for entities that are small businesses, or nonprofit private entities, as de-11 fined by the Secretary for purposes of this section. 12

13 "(d) FEE WAIVER OR REDUCTION.—The Secretary 14 shall grant a waiver from or a reduction of a fee assessed 15 under subsection (a) if the Secretary finds that the fee 16 to be paid will exceed the anticipated present and future 17 costs incurred by the Secretary in carrying out the pur-18 poses described in subsection (b) (which finding may be 19 made by the Secretary using standard costs).

20 "(e) Assessment of Fees.—

21 "(1) LIMITATION.—Fees may not be assessed
22 under subsection (a) for a fiscal year beginning after
23 the first applicable fiscal year unless the amount appropriated for salaries and expenses of the Food and
24 Drug Administration for such fiscal year is equal to

1	or greater than the amount appropriated for salaries
2	and expenses of the Food and Drug Administration
3	for the first applicable fiscal year multiplied by the
4	adjustment factor applicable to the fiscal year in-
5	volved, except that in making determinations under
6	this paragraph for the fiscal years involved there
7	shall be excluded—
8	"(A) the amounts appropriated under sub-
9	section $(f)(2)$ for the fiscal years involved; and
10	"(B) the amounts appropriated under sec-
11	tion 736(g) for such fiscal years.
12	"(2) AUTHORITY.—If under paragraph $(1)$ the
13	Secretary does not have authority to assess fees
14	under subsection (a) during a portion of a fiscal
15	year, but does at a later date in such fiscal year
16	have such authority, the Secretary, notwithstanding
17	the due date under such subsection for fees, may as-
18	sess and collect such fees at any time in such fiscal
19	year, without any modification in the rate of the
20	fees.
21	"(f) Crediting and Availability of Fees.—
22	"(1) IN GENERAL.—Fees collected for a fiscal
23	year pursuant to subsection (a) shall be credited to
24	the appropriation account for salaries and expenses
25	of the Food and Drug Administration and shall be

1	available in accordance with appropriation Acts until
2	expended without fiscal year limitation. Such sums
3	as may be necessary may be transferred from the
4	Food and Drug Administration salaries and ex-
5	penses appropriation account without fiscal year lim-
6	itation to such appropriation account for salaries
7	and expenses with such fiscal year limitation. The
8	sums transferred shall be available solely for the
9	purposes described in paragraph (1) of subsection
10	(b), and the sums are subject to allocations under
11	paragraph (2) of such subsection.
12	"(2) Authorization of appropriations.—
13	"(A) FIRST FISCAL YEAR.—For the first
14	applicable fiscal year—
15	"(i) there is authorized to be appro-
16	priated for fees under subsection (a) an
17	amount equal to the amount of increase
18	determined under subsection $(b)(1)(A)$ by
19	the Secretary (which amount shall be pub-
20	lished in the Federal Register); and
21	"(ii) in addition, there is authorized to
22	be appropriated for fees under subsection
23	(a) an amount determined by the Secretary
24	to be necessary to carry out the purpose

described in subsection $(b)(1)(B)$ (which
amount shall be so published).
"(B) SUBSEQUENT FISCAL YEARS.—For
each of the four fiscal years following the first
applicable fiscal year—
"(i) there is authorized to be appro-
priated for fees under subsection (a) an
amount equal to the amount that applied
under subparagraph (A)(i) for the first ap-
plicable fiscal year, except that such
amount shall be adjusted under paragraph
(3)(A) for the fiscal year involved; and
"(ii) in addition, there is authorized to
be appropriated for fees under subsection
(a) an amount equal to the amount that
applied under subparagraph (A)(ii) for the
first applicable fiscal year, except that such
amount shall be adjusted under paragraph
(3)(B) for the fiscal year involved.
"(3) Adjustments.—
"(A) AGENCY COST OF RESOURCES.—For
each fiscal year other than the first applicable
fiscal year, the amount that applied under para-
graph $(2)(A)(i)$ for the first applicable fiscal

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1	year shall be multiplied by the adjustment fac-
2	tor (as defined in subsection (i)).
3	"(B) RESEARCH PROGRAM.—For each fis-
4	cal year other than the first applicable fiscal
5	year, the amount that applied under paragraph
6	(2)(A)(ii) for the first applicable fiscal year
7	shall be adjusted by the Secretary (and as ad-
8	justed shall be published in the Federal Reg-
9	ister) to reflect the greater of—
10	"(i) the total percentage change that
11	occurred during the preceding fiscal year
12	in the Consumer Price Index for all urban
13	consumers (all items; U.S. city average); or
14	"(ii) the total percentage change for
15	such fiscal year in basic pay under the
16	General Schedule in accordance with sec-
17	tion 5332 of title 5, United States Code,
18	as adjusted by any locality-based com-
19	parability payment pursuant to section
20	5304 of such title for Federal employees
21	stationed in the District of Columbia.
22	"(4) Offset.—Any amount of fees collected
23	for a fiscal year under subsection (a) that exceeds
24	the amount of fees specified in appropriation Acts
25	for such fiscal year shall be credited to the appro-

priation account of the Food and Drug Administration as provided in paragraph (1), and shall be subtracted from the amount of fees that would otherwise be authorized to be collected under this section
pursuant to appropriation Acts for a subsequent fiscal year.

7 "(g) COLLECTION OF UNPAID FEES.—In any case
8 where the Secretary does not receive payment of a fee as9 sessed under subsection (a) within 30 days after it is due,
10 such fee shall be treated as a claim of the United States
11 Government subject to subchapter II of chapter 37 of title
12 31, United States Code.

13 "(h) CONSTRUCTION.—This section may not be construed as requiring that the number of full-time equivalent 14 15 positions in the Department of Health and Human Services, for officers, employers, and advisory committees not 16 17 engaged in carrying out section 409 with respect to genetic food additives be reduced to offset the number of 18 officers, employees, and advisory committees so engaged. 19 "(i) DEFINITION OF ADJUSTMENT FACTOR.—For 20 21 purposes of this section, the term 'adjustment factor' ap-22 plicable to a fiscal year is the lower of—

23 "(1) the Consumer Price Index for all urban
24 consumers (all items; United States city average) for

1	April of the preceding fiscal year divided by such
2	Index for April of the first applicable fiscal year; or
3	((2) the total of discretionary budget authority
4	provided for programs in categories other than the
5	defense category for the immediately preceding fiscal
6	year (as reported in the Office of Management and
7	Budget sequestration preview report, if available, re-
8	quired under section 254(c) of the Balanced Budget
9	and Emergency Deficit Control Act of 1985) divided
10	by such budget authority for the first applicable fis-
11	cal year (as reported in the Office of Management
12	and Budget final sequestration report submitted for
13	such year).
14	For purposes of this subsection, the terms 'budget author-
15	ity' and 'category' have the meaning given such terms in
16	the Balanced Budget and Emergency Deficit Control Act
17	of 1985.".
18	SEC. 5. EMBARGO AUTHORITY.
19	(a) Embargo.—

20 (1) TEMPORARY DETENTION.—Section
21 304(g)(1) of the Federal Food, Drug, and Cosmetic
22 Act (21 U.S.C. 334(g)(1)) is amended—

24 (i) by striking "If during" and all25 that follows through "order the device de-

(A) in the first sentence—

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1	tained" and inserting the following: "If,
2	during an inspection conducted under sec-
3	tion 704, an officer or employee of the De-
4	partment has reason to believe that a food
5	or device is in violation of this Act, such
6	officer or employee may order the food or
7	device detained"; and
8	(ii) by striking "he may authorize"
9	and inserting "the Secretary may author-
10	ize'';
11	(B) in the second and third sentences, by
12	striking "device" each place such term appears
13	and inserting "food or device";
14	(C) by striking the fourth and fifth sen-
15	tences; and
16	(D) by adding at the end the following sen-
17	tence: "A detention order under this paragraph
18	shall be considered final agency action.".
19	(2) Conforming Amendments.—Chapter III
20	of the Federal Food, Drug, and Cosmetic Act (21
21	U.S.C. 331 et seq.) is amended—
22	(A) in section $301(r)$ —
23	(i) by striking "device" the first place
24	such term appears and inserting "food or
25	device"; and

1	(ii) by striking "the device" and in-
2	serting "such food or device"; and
3	(B) in section $304(g)(2)$ , by striking "de-
4	vice" each place such term appears and insert-
5	ing "food or device".
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6 (b) DATE CERTAIN FOR PROPOSED AND FINAL 7 RULES.—Within six months of the date of the enactment 8 of this Act, the Secretary of Health and Human Services 9 shall propose a revision to the regulations in effect on such 10 date under section 304(g) of the Federal Food, Drug, and Cosmetic Act to include food. Within three months of the 11 12 date such proposed revision is published in the Federal 13 Register, the Secretary shall issue a final revision of such regulations. 14

(c) CONFIDENTIALITY.—For any food embargoed,
seized, or recalled under the Federal Food, Drug, and Cosmetic Act, the Food and Drug Administration shall disclose all necessary information without regard to business
confidentiality, if such disclosure is necessary to fully embargo, seize, or recall any adulterated food.

(d) FOOD RETAILER REGISTRATION.—All food retailers shall register with the Food and Drug Administration for the purpose of expediting recalls, embargoes, and
seizures under the Federal Food, Drug, and Cosmetic Act.

### 1 SEC. 6. RULEMAKING; EFFECTIVE DATE; PREVIOUSLY UN-2 REGULATED MARKETED ADDITIVES.

3 (a) RULEMAKING; EFFECTIVE DATE.—Not later than one year after the date of the enactment of this Act, 4 5 the Secretary of Health and Human Services shall by regulation establish criteria for carrying out section 409 of 6 7 the Federal Food, Drug, and Cosmetic Act in accordance 8 with the amendments made by section 3, and criteria for 9 carrying out section 409A of such Act (as added by section 10 4). Such amendments take effect upon the expiration of 11 the 30-day period beginning on the date on which the Sec-12 retary promulgates the final rule under the preceding sen-13 tence, subject to subsection (b).

14 (b) PREVIOUSLY UNREGULATED MARKETED ADDI-15 TIVES.—

16 (1) IN GENERAL.—In the case of a genetic food 17 additive (as defined pursuant to the amendments 18 made by section (3)) that in the United States was 19 in commercial use in food as of the day before the 20 date on which the final rule under subsection (a) is 21 promulgated, the amendments made by this Act 22 apply to the additive upon the expiration of the two-23 year period beginning on the date on which the final 24 rule is promulgated, subject to paragraph (2).

25 (2) USER FEES.—With respect to a genetic
26 food additive described in paragraph (1), such para•HR 2917 IH

graph does not waive the applicability of section
 409A of the Federal Food, Drug, and Cosmetic Act
 to a petition under section 409(b)(1) of such Act
 that is filed before the expiration of the two-year pe riod described in such paragraph.