

104TH CONGRESS
1ST SESSION

H. R.

A BILL

Referred to the Committee on

Calendar No. 29

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H. R. 956

AN ACT

To establish legal standards and procedures for product liability litigation, and for other purposes.

MARCH 15, 1995

Read the second time and placed on the calendar

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104TH CONGRESS
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IN THE SENATE OF THE UNITED STATES

MARCH 14 (legislative day, MARCH 6), 1995

Received; read the first time

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AN ACT

To establish legal standards and procedures for product liability litigation, 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 AND TABLE OF CONTENTS.**

4 (a) SHORT TITLE.—This Act may be cited as the
5 “Common Sense Product Liability and Legal Reform Act
6 of 1995”.

1 (b) TABLE OF CONTENTS.—The table of contents is
 2 as follows:

- Sec. 1. Short title and table of contents.
- Sec. 2. Findings and purposes.

TITLE I—PRODUCT LIABILITY REFORM

- Sec. 101. Applicability and preemption.
- Sec. 102. Liability rules applicable to product sellers.
- Sec. 103. Defense based on claimant’s use of intoxicating alcohol or drugs.
- Sec. 104. Misuse or alteration.
- Sec. 105. Frivolous pleadings.
- Sec. 106. Statute of repose.
- Sec. 107. Foreign products.
- Sec. 108. Definitions.

TITLE II—LIMITATION ON SPECULATIVE AND ARBITRARY DAMAGE AWARDS

- Sec. 201. Punitive damages as penalty in civil actions.
- Sec. 202. Fair share rule for noneconomic damage awards.
- Sec. 203. Limitation on noneconomic damages in health care liability actions.
- Sec. 204. Definitions.

TITLE III—BIOMATERIALS SUPPLIERS

- Sec. 301. Liability of biomaterials suppliers.
- Sec. 302. Procedures for dismissal of civil actions against biomaterials suppliers.
- Sec. 303. Definitions.

TITLE IV—LIMITATIONS ON APPLICABILITY; EFFECTIVE DATE

- Sec. 401. Application limited to interstate commerce.
- Sec. 402. Effect on other law.
- Sec. 403. Federal cause of action precluded.
- Sec. 404. Effective date.

3 **SEC. 2. FINDINGS AND PURPOSES.**

4 (a) FINDINGS.—The Congress finds that—

- 5 (1) the civil justice system, which is designed to
- 6 safeguard our most cherished rights, to remedy in-
- 7 justices, and to defend our liberty, is increasingly
- 8 being deployed to abridge our rights, create injus-
- 9 tice, and destroy our liberty;

1 (2) our Nation is overly litigious, the civil jus-
2 tice system is overcrowded, sluggish, and excessively
3 costly, and the costs of lawsuits, both direct and in-
4 direct, are inflicting serious and unnecessary injury
5 on the national economy;

6 (3) excessive, unpredictable, and often arbitrary
7 damage awards and unfair allocations of liability
8 have a direct and undesirable effect on interstate
9 commerce by increasing the cost and decreasing the
10 availability of goods and services;

11 (4) the rules of law governing product liability
12 actions, damage awards, and allocations of liability
13 have evolved inconsistently within and among the
14 several States, resulting in a complex, contradictory,
15 and uncertain regime that is inequitable to both
16 plaintiffs and defendants and unduly burdens inter-
17 state commerce;

18 (5) as a result of excessive, unpredictable, and
19 often arbitrary damage awards and unfair alloca-
20 tions of liability, consumers have been adversely af-
21 fected through the withdrawal of products, produc-
22 ers, services, and service providers from the national
23 market, and from excessive liability costs passed on
24 to them through higher prices;

1 (6) excessive, unpredictable, and often arbitrary
2 damage awards and unfair allocations of liability
3 jeopardize the financial well-being of many individ-
4 uals as well as entire industries, particularly the Na-
5 tion's small businesses, and adversely affects govern-
6 ments, taxpayers, nonprofit entities and volunteer
7 organizations;

8 (7) the excessive costs of the civil justice system
9 undermine the ability of American companies to
10 compete internationally, and serve to decrease the
11 number of jobs and the amount of productive capital
12 in the national economy;

13 (8) the unpredictability of damage awards is in-
14 equitable to both plaintiffs and defendants and has
15 added considerably to the high cost of liability insur-
16 ance, making it difficult for producers, consumers,
17 and individuals to protect their liability with any de-
18 gree of confidence and at a reasonable cost;

19 (9) because of the national scope of the prob-
20 lems created by the defects in the civil justice sys-
21 tem, it is not possible for the several States to enact
22 laws that fully and effectively respond to those prob-
23 lems;

1 (10) it is the constitutional role of the national
2 government to remove barriers to interstate com-
3 merce; and

4 (11) there is a need to restore rationality, cer-
5 tainty, and fairness to the civil justice system in
6 order to protect against excessive, arbitrary, and un-
7 certain damage awards and to reduce the volume,
8 costs, and delay of litigation.

9 (b) PURPOSES.—Based upon the powers contained in
10 Article I, Section 8, Clause 3 of the United States Con-
11 stitution, the purposes of this Act are to promote the free
12 flow of goods and services and to lessen burdens on inter-
13 state commerce by—

14 (1) establishing certain uniform legal principles
15 of product liability which provide a fair balance
16 among the interests of product users, manufactur-
17 ers, and product sellers;

18 (2) placing reasonable limits on damages over
19 and above the actual damages suffered by a claim-
20 ant;

21 (3) ensuring the fair allocation of liability in
22 civil actions;

23 (4) reducing the unacceptable costs and delays
24 of our civil justice system caused by excessive litiga-
25 tion which harm both plaintiffs and defendants; and

1 (5) establishing greater fairness, rationality,
2 and predictability in the civil justice system.

3 **TITLE I—PRODUCT LIABILITY**
4 **REFORM**

5 **SEC. 101. APPLICABILITY AND PREEMPTION.**

6 (a) PREEMPTION.—This title governs any product li-
7 ability action brought in any State or Federal court, on
8 any theory for harm caused by a product. A civil action
9 brought for commercial loss shall be governed only by ap-
10 plicable commercial or contract law.

11 (b) RELATIONSHIP TO STATE LAW.—This title su-
12 persedes State law only to the extent that State law ap-
13 plies to an issue covered by this title. Any issue that is
14 not governed by this title shall be governed by otherwise
15 applicable State or Federal law.

16 **SEC. 102. LIABILITY RULES APPLICABLE TO PRODUCT**
17 **SELLERS.**

18 (a) GENERAL RULE.—Except as provided in sub-
19 section (b), in any product liability action, a product seller
20 other than a manufacturer shall be liable to a claimant
21 for harm only if the claimant establishes that—

22 (1)(A) the product which allegedly caused the
23 harm complained of was sold by the product seller;
24 (B) the product seller failed to exercise reasonable
25 care with respect to the product; and (C) such fail-

1 ure to exercise reasonable care was a proximate
2 cause of the claimant's harm; or

3 (2)(A) the product seller made an express war-
4 ranty applicable to the product which allegedly
5 caused the harm complained of, independent of any
6 express warranty made by a manufacturer as to the
7 same product; (B) the product failed to conform to
8 the warranty; and (C) the failure of the product to
9 conform to the warranty caused the claimant's
10 harm; or

11 (3) the product seller engaged in intentional
12 wrongdoing as determined under applicable State
13 law and such intentional wrongdoing was a proxi-
14 mate cause of the harm complained of by the claim-
15 ant.

16 For purposes of paragraph (1)(B), a product seller shall
17 not be considered to have failed to exercise reasonable care
18 with respect to the product based upon an alleged failure
19 to inspect a product where there was no reasonable oppor-
20 tunity to inspect the product in a manner which would,
21 in the exercise of reasonable care, have revealed the aspect
22 of the product which allegedly caused the claimant's harm.

23 (b) EXCEPTION.—In a product liability action, a
24 product seller shall be liable for harm to the claimant

1 caused by such product as if the product seller were the
2 manufacturer of such product if—

3 (1) the manufacturer is not subject to service of
4 process under the laws of any State in which the ac-
5 tion might have been brought; or

6 (2) the court determines that the claimant
7 would be unable to enforce a judgment against the
8 manufacturer.

9 (c) RENTAL AND LEASES.—Notwithstanding any
10 other provision of law, any person, except a person ex-
11 cluded from the definition of product seller, engaged in
12 the business of renting or leasing a product shall be sub-
13 ject to liability pursuant to subsection (a) of this section,
14 but shall not be liable to a claimant for the tortious act
15 of another solely by reason of ownership of such product.

16 **SEC. 103. DEFENSE BASED ON CLAIMANT'S USE OF INTOXI-**
17 **CATING ALCOHOL OR DRUGS.**

18 (a) GENERAL RULE.—In any product liability action,
19 it shall be a complete defense to such action if—

20 (1) the claimant was intoxicated or was under
21 the influence of intoxicating alcohol or any drug
22 when the accident or other event which resulted in
23 such claimant's harm occurred; and

1 (2) the claimant, as a result of the influence of
2 the alcohol or drug, was more than 50 percent re-
3 sponsible for such accident or other event.

4 (b) CONSTRUCTION.—For purposes of subsection
5 (a)—

6 (1) the determination of whether a person was
7 intoxicated or was under the influence of intoxicat-
8 ing alcohol or any drug shall be made pursuant to
9 applicable State law; and

10 (2) the term “drug” means any controlled sub-
11 stance as defined in the Controlled Substances Act
12 (21 U.S.C. 802(6)) that has been taken by the
13 claimant other than in accordance with the terms of
14 a lawfully issued prescription.

15 **SEC. 104. MISUSE OR ALTERATION.**

16 (a) GENERAL RULE.—In a product liability action,
17 the damages for which a defendant is otherwise liable
18 under State law shall be reduced by the percentage of re-
19 sponsibility for the claimant’s harm attributable to misuse
20 or alteration of a product by any person if the defendant
21 establishes by a preponderance of the evidence that such
22 percentage of the claimant’s harm was proximately caused
23 by—

24 (1) a use or alteration of a product in violation
25 of, or contrary to, a defendant’s express warnings or

1 instructions if the warnings or instructions are ade-
2 quate as determined pursuant to applicable State
3 law, or

4 (2) a use or alteration of a product involving a
5 risk of harm which was known or should have been
6 known by the ordinary person who uses or consumes
7 the product with the knowledge common to the class
8 of persons who used or would be reasonably antici-
9 pated to use the product.

10 (b) **WORKPLACE INJURY.**—Notwithstanding sub-
11 section (a), the damage for which a defendant is otherwise
12 liable under State law shall not be reduced by the percent-
13 age of responsibility for the claimant’s harm attributable
14 to misuse or alteration of the product by the claimant’s
15 employer or any co-employee who is immune from suit by
16 the claimant pursuant to the State law applicable to work-
17 place injuries.

18 **SEC. 105. FRIVOLOUS PLEADINGS.**

19 (a) **GENERAL RULE.**—

20 (1) **SIGNING OF PLEADING.**—The signing or
21 verification of a pleading in a product liability action
22 in a State court subject to this title constitutes a
23 certificate that to the signatory’s or verifier’s best
24 knowledge, information, and belief, formed after rea-

1 sonable inquiry, the pleading is not frivolous as de-
2 termined under paragraph (2).

3 (2) DEFINITIONS.—

4 (A) For purposes of this section, a plead-
5 ing is frivolous if the pleading is—

6 (i) groundless and brought in bad
7 faith;

8 (ii) groundless and brought for the
9 purpose of harassment; or

10 (iii) groundless and interposed for any
11 improper purpose, such as to cause unnec-
12 essary delay or needless increase in the
13 cost of litigation.

14 (B) For purposes of subparagraph (A), the
15 term “groundless” means—

16 (i) no basis in fact; or

17 (ii) not warranted by existing law or
18 a good faith argument for the extension,
19 modification, or reversal of existing law.

20 (b) DETERMINATION THAT PLEADING FRIVO-
21 LOUS.—

22 (1) MOTION FOR DETERMINATION.—Not later
23 than 60 days after the date a pleading in a product
24 liability action in a State court is filed, a party to

1 the action may make a motion that the court deter-
2 mine if the pleading is frivolous.

3 (2) COURT ACTION.—The court in a product li-
4 ability action in a State court shall on the motion
5 of a party or on its own motion determine if a plead-
6 ing is frivolous.

7 (c) CONSIDERATIONS.—In making its determination
8 of whether a pleading is frivolous, the court shall take into
9 account—

10 (1) the multiplicity of parties;

11 (2) the complexity of the claims and defenses;

12 (3) the length of time available to the party to
13 investigate and conduct discovery; and

14 (4) affidavits, depositions, and any other rel-
15 evant matter.

16 (d) SANCTION.—If the court determines that a plead-
17 ing is frivolous, the court shall impose an appropriate
18 sanction on the signatory or verifier of the pleading. The
19 sanction may include one or more of the following:

20 (1) the striking of a pleading or the offending
21 portion thereof;

22 (2) the dismissal of a party; or

23 (3) an order to pay to a party who stands in
24 opposition to the offending pleading the amounts of
25 the reasonable expenses incurred because of the fil-

1 ing of the pleading, including costs, reasonable at-
2 torney's fees, witness fees, fees of experts, and depo-
3 sition expenses.

4 (e) CONSTRUCTION.—For purposes of this section—

5 (1) a general denial does not constitute a frivo-
6 lous pleading; and

7 (2) the amount requested for damages does not
8 constitute a frivolous pleading.

9 **SEC. 106. STATUTE OF REPOSE.**

10 (a) GENERAL RULE.—A product liability action shall
11 be barred unless the complaint is served and filed within
12 15 years of the date of delivery of the product to its first
13 purchaser or lessee, who was not engaged in the business
14 of selling or leasing the product or of using the product
15 as a component in the manufacture of another product.

16 (b) EXCEPTION.—Subsection (a)—

17 (1) does not bar a product liability action
18 against a defendant who made an express warranty
19 in writing as to the safety of the specific product in-
20 volved which was longer than 15 years, but it will
21 apply at the expiration of such warranty,

22 (2) does not apply to a physical illness the evi-
23 dence of which does not ordinarily appear less than
24 15 years after the first exposure to the product, and

1 (3) does not affect the limitations period estab-
2 lished by the General Aviation Revitalization Act of
3 1994.

4 **SEC. 107. FOREIGN PRODUCTS.**

5 (a) GENERAL RULE.—In any product liability action
6 for injury that was sustained in the United States and
7 that relates to the purchase or use of a product manufac-
8 tured outside the United States by a foreign manufac-
9 turer, the Federal court in which such action is brought
10 shall have jurisdiction over such manufacturer if the man-
11 ufacturer knew or reasonably should have known that the
12 product would be imported for sale or use in the United
13 States.

14 (b) ADMISSION.—If in any product liability action a
15 foreign manufacturer of the product involved in such ac-
16 tion fails to furnish any testimony, document, or other
17 thing upon a duly issued discovery order by the court in
18 such action, such failure shall be deemed an admission of
19 any fact with respect to which the discovery order relates.

20 (c) PROCESS.—Process in an action described in sub-
21 section (a) may be served wherever the foreign manufac-
22 turer is located, has an agent, or transacts business.

23 **SEC. 108. DEFINITIONS.**

24 As used in this title:

1 (1) The term “claimant” means any person who
2 brings a product liability action and any person on
3 whose behalf such an action is brought. If such an
4 action is brought through or on behalf of an estate,
5 the term includes the claimant’s decedent. If such
6 action is brought through or on behalf of a minor
7 or incompetent, the term includes the claimant’s
8 legal guardian.

9 (2) The term “commercial loss” means any loss
10 of or damage to a product itself incurred in the
11 course of the ongoing business enterprise consisting
12 of providing goods or services for compensation.

13 (3) The term “economic loss” means any pecu-
14 niary loss resulting from harm (including the loss of
15 earnings, medical expense loss, replacement services
16 loss, loss due to death, and burial costs) to the ex-
17 tent recovery for such loss is allowed under applica-
18 ble State law.

19 (4) The term “harm” means any physical in-
20 jury, illness, disease, or death or damage to property
21 caused by a product. The term does not include
22 commercial loss or loss or damage to a product it-
23 self.

24 (5) The term “manufacturer” means—

1 (A) any person who is engaged in a busi-
2 ness to produce, create, make, or construct any
3 product (or component part of a product) and
4 who (i) designs or formulates the product (or
5 component part of the product), (ii) has en-
6 gaged another person to design or formulate
7 the product (or component part of the product),
8 or (iii) uses the design or formulation of the
9 product developed by another person;

10 (B) a product seller of the product who,
11 before placing the product in the stream of
12 commerce—

13 (i) designs or formulates or has en-
14 gaged another person to design or formu-
15 late an aspect of the product after the
16 product was initially made by another, or

17 (ii) produces, creates, makes, or con-
18 structs such aspect of the product, or

19 (C) any product seller not described in
20 subparagraph (B) which holds itself out as a
21 manufacturer to the user of the product.

22 (6) The term “noneconomic loss” means subjec-
23 tive, nonmonetary loss resulting from harm, includ-
24 ing pain, suffering, inconvenience, mental suffering,
25 emotional distress, loss of society and companion-

1 ship, loss of consortium, injury to reputation, and
2 humiliation.

3 (7) The term “person” means any individual,
4 corporation, company, association, firm, partnership,
5 society, joint stock company, or any other entity (in-
6 cluding any governmental entity).

7 (8)(A) The term “product” means any object,
8 substance, mixture, or raw material in a gaseous,
9 liquid, or solid state which—

10 (i) is capable of delivery itself or as an as-
11 sembled whole, in a mixed or combined state, or
12 as a component part or ingredient;

13 (ii) is produced for introduction into trade
14 or commerce;

15 (iii) has intrinsic economic value; and

16 (iv) is intended for sale or lease to persons
17 for commercial or personal use.

18 (B) The term does not include—

19 (i) human tissue, human organs, human
20 blood, and human blood products; or

21 (ii) electricity, water delivered by a utility,
22 natural gas, or steam.

23 (9) The term “product liability action” means
24 a civil action brought on any theory for harm caused
25 by a product or product use.

1 (10) The term “product seller” means a person
2 who, in the course of a business conducted for that
3 purpose, sells, distributes, rents, leases, prepares,
4 blends, packages, labels a product, is otherwise in-
5 volved in placing a product in the stream of com-
6 merce, or installs, repairs, or maintains the harm-
7 causing aspect of a product. The term does not in-
8 clude—

9 (A) a seller or lessor of real property;

10 (B) a provider of professional services in
11 any case in which the sale or use of a product
12 is incidental to the transaction and the essence
13 of the transaction is the furnishing of judg-
14 ment, skill, or services; or

15 (C) any person who—

16 (i) acts in only a financial capacity
17 with respect to the sale of a product; or

18 (ii) leases a product under a lease ar-
19 rangement in which the selection, posses-
20 sion, maintenance, and operation of the
21 product are controlled by a person other
22 than the lessor.

23 (11) The term “State” means any State of the
24 United States, the District of Columbia, Common-
25 wealth of Puerto Rico, the Northern Mariana Is-

1 lands, the Virgin Islands, Guam, American Samoa,
2 and any other territory or possession of the United
3 States, or any political subdivision of any of the
4 foregoing.

5 **TITLE II—LIMITATION ON SPEC-**
6 **ULATIVE AND ARBITRARY**
7 **DAMAGE AWARDS**

8 **SEC. 201. PUNITIVE DAMAGES AS PENALTY IN CIVIL AC-**
9 **TIONS.**

10 (a) GENERAL RULE.—Punitive damages may, to the
11 extent permitted by applicable State law, be awarded in
12 any civil action for harm in any Federal or State court
13 against a defendant if the claimant establishes by clear
14 and convincing evidence that the harm suffered was result
15 of conduct—

16 (1) specifically intended to cause harm, or

17 (2) conduct manifesting a conscious, flagrant
18 indifference to the rights or safety of others.

19 (b) PROPORTIONAL AWARDS.—The amount of puni-
20 tive damages that may be awarded in any civil action sub-
21 ject to this title shall not exceed 3 times the amount of
22 damages awarded to the claimant for economic loss, or
23 \$250,000, whichever is greater. This section shall be ap-
24 plied by the court and shall not be disclosed to the jury.

1 (c) APPLICABILITY.—Except as provided in section
2 401, this section shall apply to any civil action brought
3 in any Federal or State court on any theory where punitive
4 damages are sought. This section does not create a cause
5 of action for punitive damages. This section does not pre-
6 empt or supersede any State or Federal law to the extent
7 that such law would further limit the award of punitive
8 damages.

9 (d) BIFURCATION.—At the request of any party, the
10 trier of fact shall consider in a separate proceeding wheth-
11 er punitive damages are to be awarded and the amount
12 of such award. If a separate proceeding is requested, evi-
13 dence relevant only to the claim of punitive damages, as
14 determined by applicable State law, shall be inadmissible
15 in any proceeding to determine whether compensatory
16 damages are to be awarded.

17 (e) CONSIDERATIONS.—In determining the amount
18 of punitive damages, the trier of fact shall consider all rel-
19 evant, admissible evidence, including—

20 (1) the severity of the harm caused by the con-
21 duct of the defendant,

22 (2) the duration of the conduct or any conceal-
23 ment of it by the defendant,

24 (3) the profitability of the specific conduct that
25 caused the harm to the defendant,

1 (4) the number of products sold, the frequency
2 of services provided, or the type of activities con-
3 ducted by the defendant of the kind causing the
4 harm complained of by the claimant,

5 (5) awards of punitive damages to persons simi-
6 larly situated to the claimant,

7 (6) possibility of prospective awards of compen-
8 satory damages to persons similarly situated to the
9 claimant,

10 (7) any criminal penalties imposed on the de-
11 fendant as a result of the conduct complained of by
12 the claimant,

13 (8) the amount of any civil and administrative
14 fines and penalties assessed against the defendant as
15 a result of the conduct complained of by the claim-
16 ant, and

17 (9) whether the foregoing considerations have
18 been a factor in any prior proceeding involving the
19 defendant.

20 (f) DRUGS AND DEVICES.—

21 (1)(A) Punitive damages shall not be awarded
22 against a manufacturer or product seller of a drug
23 (as defined in section 201(g)(1) of the Federal
24 Food, Drug, and Cosmetic Act (21 U.S.C.
25 321(g)(1)) or medical device (as defined in section

1 201(h) of the Federal Food, Drug, and Cosmetic
2 Act (21 U.S.C. 321(h)) which caused the claimant's
3 harm where—

4 (i) such drug or device was subject to pre-
5 market approval by the Food and Drug Admin-
6 istration with respect to the safety of the for-
7 mulation or performance of the aspect of such
8 drug or device which caused the claimant's
9 harm or the adequacy of the packaging or label-
10 ing of such drug or device, and such drug was
11 approved by the Food and Drug Administra-
12 tion; or

13 (ii) the drug is generally recognized as safe
14 and effective pursuant to conditions established
15 by the Food and Drug Administration and ap-
16 plicable regulations, including packaging and la-
17 beling regulations.

18 (B) Subparagraph (A) shall not apply in any
19 case in which the defendant, before or after pre-
20 market approval of a drug or device—

21 (i) intentionally and wrongfully withheld
22 from or misrepresented to the Food and Drug
23 Administration information concerning such
24 drug or device required to be submitted under
25 the Federal Food, Drug, and Cosmetic Act (21

1 U.S.C. 301 et seq.) or section 351 of the Public
2 Health Service Act (42 U.S.C. 262) that is ma-
3 terial and relevant to the harm suffered by the
4 claimant, or

5 (ii) made an illegal payment to an official
6 or employee of the Food and Drug Administra-
7 tion for the purpose of securing or maintaining
8 approval of such drug or device.

9 (2) PACKAGING.—In a product liability action
10 for harm which is alleged to relate to the adequacy
11 of the packaging (or labeling relating to such pack-
12 aging) of a drug which is required to have tamper-
13 resistant packaging under regulations of the Sec-
14 retary of Health and Human Services (including la-
15 beling regulations related to such packaging), the
16 manufacturer of the drug shall not be held liable for
17 punitive damages unless the drug is found by the
18 court by clear and convincing evidence to be sub-
19 stantially out of compliance with such regulations.

20 **SEC. 202. FAIR SHARE RULE FOR NONECONOMIC DAMAGE**
21 **AWARDS.**

22 (a) FAIR SHARE OF LIABILITY IMPOSED ACCORDING
23 TO SHARE OF FAULT.—In any product liability or other
24 civil action brought in State or Federal court, a defendant
25 shall be liable only for the amount of noneconomic dam-

1 ages attributable to such defendant in direct proportion
2 to such defendant's share of fault or responsibility for the
3 claimant's actual damages, as determined by the trier of
4 fact. In all such cases, the liability of a defendant for non-
5 economic damages shall be several and not joint.

6 (b) APPLICABILITY.—Except as provided in section
7 402, this section shall apply to any product liability or
8 other civil action brought in any Federal or State court
9 on any theory where noneconomic damages are sought.
10 This section does not preempt or supersede any State or
11 Federal law to the extent that such law would further limit
12 the application of the theory of joint liability to any kind
13 of damages.

14 **SEC. 203. LIMITATION ON NONECONOMIC DAMAGES IN**
15 **HEALTH CARE LIABILITY ACTIONS.**

16 (a) MAXIMUM AWARD OF NONECONOMIC DAM-
17 AGES.—In any health care liability action, in addition to
18 actual damages or punitive damages, or both, a claimant
19 may also be awarded noneconomic damages, including
20 damages awarded to compensate injured feelings, such as
21 pain and suffering and emotional distress. The maximum
22 amount of such damages that may be awarded to a claim-
23 ant shall be \$250,000. Such maximum amount shall apply
24 regardless of the number of parties against whom the ac-
25 tion is brought, and regardless of the number of claims

1 or actions brought with respect to the health care injury.
2 An award for future noneconomic damages shall not be
3 discounted to present value. The jury shall not be in-
4 formed about the limitation on noneconomic damages, but
5 an award for noneconomic damages in excess of \$250,000
6 shall be reduced either before the entry of judgment or
7 by amendment of the judgment after entry. An award of
8 damages for noneconomic losses in excess of \$250,000
9 shall be reduced to \$250,000 before accounting for any
10 other reduction in damages required by law. If separate
11 awards of damages for past and future noneconomic dam-
12 ages are rendered and the combined award exceeds
13 \$250,000, the award of damages for future noneconomic
14 losses shall be reduced first.

15 (b) APPLICABILITY.—Except as provided in section
16 401, this section shall apply to any health care liability
17 action brought in any Federal or State court on any theory
18 or pursuant to any alternative dispute resolution process
19 where noneconomic damages are sought. This section does
20 not create a cause of action for noneconomic damages.
21 This section does not preempt or supersede any State or
22 Federal law to the extent that such law would further limit
23 the award of noneconomic damages. This section does not
24 preempt any State law enacted before the date of the en-

1 actment of this Act that places a cap on the total liability
2 in a health care liability action.

3 (c) DEFINITIONS.—As used in this section:

4 (1) The term “claimant” means any person who
5 asserts a health care liability claim or brings a
6 health care liability action, including a person who
7 asserts or claims a right to legal or equitable con-
8 tribution, indemnity or subrogation, arising out of a
9 health care liability claim or action, and any person
10 on whose behalf such a claim is asserted or such an
11 action is brought, whether deceased, incompetent or
12 a minor.

13 (2) The term “economic loss” has the same
14 meaning as defined at section 204(4).

15 (3) The term “health care liability action”
16 means a civil action brought in a State or Federal
17 court or pursuant to any alternative dispute resolu-
18 tion process, against a health care provider, an en-
19 tity which is obligated to provide or pay for health
20 benefits under any health plan (including any person
21 or entity acting under a contract or arrangement to
22 provide or administer any health benefit), or the
23 manufacturer, distributor, supplier, marketer, pro-
24 moter, or seller of a medical product, in which the
25 claimant alleges a claim (including third party

1 claims, cross claims, counter claims, or distribution
2 claims) based upon the provision of (or the failure
3 to provide or pay for) health care services or the use
4 of a medical product, regardless of the theory of li-
5 ability on which the claim is based, or the number
6 of plaintiffs, or defendants or causes of action.

7 **SEC. 204. DEFINITIONS.**

8 As used in this title:

9 (1) The term “actual damages” means damages
10 awarded to pay for economic loss.

11 (2) The term “claimant” means any person who
12 brings a civil action and any person on whose behalf
13 such an action is brought. If such action is brought
14 through or on behalf of an estate, the term includes
15 the claimant’s decedent. If such action is brought
16 through or on behalf of a minor or incompetent, the
17 term includes the claimant’s legal guardian.

18 (3) The term “clear and convincing evidence” is
19 that measure or degree of proof that will produce in
20 the mind of the trier of fact a firm belief or convic-
21 tion as to the truth of the allegations sought to be
22 established. The level of proof required to satisfy
23 such standard is more than that required under pre-
24 ponderance of the evidence, but less than that re-
25 quired for proof beyond a reasonable doubt.

1 (4) The term “economic loss” means any pecu-
2 niary loss resulting from harm (including the loss of
3 earnings, medical expense loss, replacement services
4 loss, loss due to death, and burial costs), to the ex-
5 tent recovery for such loss is allowed under applica-
6 ble State law.

7 (5) The term “harm” means any legally cog-
8 nizable wrong or injury for which punitive damages
9 may be imposed.

10 (6) The term “noneconomic damages” means
11 damages other than punitive damages or actual
12 damages.

13 (7) The term “punitive damages” means dam-
14 ages awarded against any person or entity to punish
15 or deter such person or entity, or others, from en-
16 gaging in similar behavior in the future.

17 (8) The term “State” means any State of the
18 United States, the District of Columbia, Common-
19 wealth of Puerto Rico, the Northern Mariana Is-
20 lands, the Virgin Islands, Guam, American Samoa,
21 and any other territory or possession of the United
22 States, or any political subdivision of any of the
23 foregoing.

1 **TITLE III—BIOMATERIALS**
2 **SUPPLIERS**

3 **SEC. 301. LIABILITY OF BIOMATERIALS SUPPLIERS.**

4 A biomaterials supplier may, to the extent required
5 and permitted by any other applicable law, be liable for
6 harm to a claimant caused by a medical device, only if
7 the claimant in a product liability action shows that the
8 conduct of the biomaterials supplier was an actual and
9 proximate cause of the harm to the claimant and—

10 (1) the raw materials or component parts deliv-
11 ered by the biomaterials supplier either—

12 (A) did not constitute the product de-
13 scribed in the contract between the biomaterials
14 supplier and the person who contracted for de-
15 livery of the product; or

16 (B) failed to meet any specifications that
17 were—

18 (i) provided to the biomaterials sup-
19 plier and not expressly repudiated by the
20 biomaterials supplier prior to acceptance of
21 delivery of the raw materials or component
22 parts;

23 (ii)(I) provided to the biomaterials
24 supplier;

1 (II) provided to the manufacturer by
2 the biomaterials supplier; or

3 (III) contained in a master file that
4 was submitted by the biomaterials supplier
5 to the Secretary of Health and Human
6 Services and that is currently maintained
7 by the biomaterials supplier of purposes of
8 premarket approval of medical devices; or

9 (iii)(I) included in the submissions for
10 the purposes of premarket approval or re-
11 view by the Secretary of Health and
12 Human Services under section 510, 513,
13 515, or 520 of the Federal Food, Drug,
14 and Cosmetic Act (21 U.S.C. 360, 360c,
15 360e, or 360j); and

16 (II) have received clearance from the
17 Secretary of Health and Human Services,
18 if such specifications were provided by the
19 manufacturer to the biomaterials supplier
20 and were not expressly repudiated by the
21 biomaterials supplier prior to the accept-
22 ance by the raw materials or component
23 parts;

24 (2) the biomaterials supplier intentionally and
25 wrongfully withheld or misrepresented information

1 that is material and relevant to the harm suffered
2 by the claimant; or

3 (3) the biomaterials supplier had actual knowl-
4 edge of prospective fraudulent or malicious activities
5 in the use of its supplies where such activities are
6 relevant to the harm suffered by the claimant.

7 **SEC. 302. PROCEDURES FOR DISMISSAL OF CIVIL ACTIONS**

8 **AGAINST BIOMATERIALS SUPPLIERS.**

9 (a) MOTION TO DISMISS.—

10 (1) GENERAL RULE.—Any biomaterials supplier
11 who is a defendant in any product liability action in-
12 volving a medical device which allegedly caused the
13 harm for which the action is brought and who did
14 not take part in the design, manufacture, or sale of
15 such medical device may, at any time during which
16 a motion to dismiss may be filed under an applicable
17 law, move to dismiss the action on the grounds
18 that—

19 (A) the claimant has failed to establish
20 that the supplier furnished raw materials or
21 component parts in violation of applicable con-
22 tractual requirements or specifications agreed
23 to by the biomaterials supplier; or

24 (B) the claimant has failed to comply with
25 the requirements of subsection (b).

1 (2) EXCEPTION.—The biomaterials supplier
2 may not move to dismiss the action if—

3 (A) the biomaterials supplier intentionally
4 and wrongfully withheld or misrepresented in-
5 formation that is material and relevant to the
6 harm suffered by the claimant; or

7 (B) the biomaterials supplier had actual
8 knowledge of prospective fraudulent or mali-
9 cious activities in the use of its supplies where
10 such activities are relevant to the harm suffered
11 by the claimant.

12 (b) MANUFACTURER OF MEDICAL DEVICE SHALL BE
13 NAMED A PARTY.—The claimant shall be required to
14 name the manufacturer of the medical device to which the
15 biomaterials supplier furnished raw materials or compo-
16 nent parts as a party to the product liability action, un-
17 less—

18 (1) the manufacturer is subject to service of
19 process solely in a jurisdiction in which the
20 biomaterials supplier is not domiciled or subject to
21 a service of process; or

22 (2) an action against the manufacturer is
23 barred by applicable law.

1 (c) PROCEEDINGS ON MOTION TO DISMISS.—The fol-
2 lowing rules shall apply to any proceeding on a motion
3 to dismiss filed under this section:

4 (1) AFFIDAVITS RELATING TO STATUS OF DE-
5 FENDANT.—

6 (A) DEFENDANT AFFIDAVIT.—The defend-
7 ant in the action may support a motion to dis-
8 miss by filing an affidavit demonstrating that
9 defendant is a biomaterials supplier and that it
10 is neither the manufacturer nor the product
11 seller of the medical device which caused the
12 harm alleged by the claimant.

13 (B) RESPONSE TO MOTION TO DISMISS.—
14 In response to a motion to dismiss described in
15 this section, the claimant may submit an affida-
16 vit demonstrating why it asserts that—

17 (i) the defendant who filed the motion
18 to dismiss is not a biomaterials supplier
19 with respect to the medical device which
20 caused the harm alleged by the claimant;

21 (ii) on what basis it asserts that the
22 supplier furnished raw materials or compo-
23 nent parts in violation of applicable con-
24 tractual requirements or specifications
25 agreed to by the biomaterials supplier;

1 (iii) the biomaterials supplier inten-
2 tionally and wrongfully withheld or mis-
3 represented information that is material
4 and relevant to the harm suffered by the
5 claimant; or

6 (iv) the biomaterials supplier had ac-
7 tual knowledge of prospective fraudulent or
8 malicious activities in the use of its sup-
9 plies where such activities are relevant to
10 the harm suffered by the claimant.

11 (2) EFFECT OF MOTION TO DISMISS ON DIS-
12 COVERY.—If a defendant files a motion to dismiss,
13 no discovery shall be permitted in connection with
14 the action that is the subject of the motion, unless
15 the affidavits submitted in accordance with this
16 section raise material issues of fact concerning
17 whether—

18 (A) the supplier furnished raw materials or
19 component parts in violation of applicable con-
20 tractual requirements or specifications agreed
21 to by the biomaterials supplier;

22 (B) the biomaterials supplier intentionally
23 and wrongfully withheld or misrepresented in-
24 formation that is material and relevant to the
25 harm suffered by the claimant; or

1 (C) the biomaterials supplier had actual
2 knowledge of prospective fraudulent or mali-
3 cious activities in the use of its supplies where
4 such activities are relevant to the harm suffered
5 by the claimant.

6 Any such discovery shall be limited solely to such
7 material facts.

8 (3) RESPONSE TO MOTION TO DISMISS.—The
9 court shall rule on the motion to dismiss solely on
10 the basis of the affidavits filed under this section
11 and on the basis of any evidence developed in the
12 course of discovery under paragraph (2) and subse-
13 quently submitted to the court in accordance with
14 applicable rules of evidence.

15 (d) ATTORNEY FEES.—The court shall require the
16 claimant to compensate the biomaterials supplier for at-
17 torney fees and costs, if—

18 (1) the claimant named or joined the
19 biomaterials supplier; and

20 (2) the court found the claim against the
21 biomaterials supplier to be without merit and frivo-
22 lous.

23 **SEC. 303. DEFINITIONS.**

24 For purposes of this title:

1 (1) The term “biomaterials supplier” means an
2 entity that directly or indirectly supplies, or licenses
3 another person to supply, a component part or raw
4 material for use in the manufacture of a medical de-
5 vice—

6 (A) that is intended by the manufacturer
7 of the device—

8 (i) to be placed into a surgically or
9 naturally formed or existing cavity of the
10 body for a period of at least 30 days; or

11 (ii) to remain in contact with bodily
12 fluids of internal human tissue through a
13 surgically produced opening for a period of
14 less than 30 days; and

15 (B) suture materials used in implant pro-
16 cedures.

17 (2) Notwithstanding paragraph (1), the term
18 “biomaterials supplier” excludes any person, with re-
19 spect to a medical device which is the subject of a
20 product liability action—

21 (A) who is engaged in the manufacture,
22 preparation, propagation, compounding, or
23 processing (as defined in section 510(a)(1) of
24 the Federal Food, Drug, and Cosmetic Act (21
25 U.S.C. 360(a)(1)) of the medical device, and

1 has or should have registered with the Sec-
2 retary of Health and Human Services pursuant
3 to section 510 of the Federal Food, Drug, and
4 Cosmetic Act (21 U.S.C. 360) and the regula-
5 tions issued under such section, and has or
6 should have included the medical device on a
7 list of devices filed with the Secretary of Health
8 and Human Services pursuant to section 510(j)
9 of such Act (21 U.S.C. 360(j)) and the regula-
10 tions issued under such section; or

11 (B) who, in the course of a business con-
12 ducted for that purpose, has sold, distributed,
13 leased, packaged, labeled, or otherwise placed
14 the implant in the stream of commerce after it
15 was manufactured.

16 (3) The term “harm” means any physical in-
17 jury, illness, disease, or death or damage to property
18 caused by a product. The term does not include
19 commercial loss or loss or damage to a product
20 itself.

21 (4) The term “product liability action” means
22 a civil action brought on any theory for harm caused
23 by a product or product use.

1 **TITLE IV—LIMITATIONS ON AP-**
2 **PLICABILITY; EFFECTIVE**
3 **DATE**

4 **SEC. 401. APPLICATION LIMITED TO INTERSTATE COM-**
5 **MERCE.**

6 Titles I, II, and III shall apply only to product liabil-
7 ity or other civil actions affecting interstate commerce.
8 For purposes of the preceding sentence, the term “inter-
9 state commerce” means commerce among the several
10 States or with foreign nations, or in any territory of the
11 United States or in the District of Columbia, or between
12 any such territory and another, or between any such terri-
13 tory and any State or foreign nation, or between the Dis-
14 trict of Columbia and any State or territory or foreign
15 nation.

16 **SEC. 402. EFFECT ON OTHER LAW.**

17 Nothing in title I, II, or III shall be construed to—

18 (1) waive or affect any defense of sovereign im-
19 munity asserted by any State under any law;

20 (2) supersede any Federal law;

21 (3) waive or affect any defense of sovereign im-
22 munity asserted by the United States;

23 (4) affect the applicability of any provision of
24 chapter 97 of title 28, United States Code;

1 (5) preempt State choice-of-law rules with re-
2 spect to claims brought by a foreign nation or a citi-
3 zen of a foreign nation; or

4 (6) affect the right of any court to transfer
5 venue or to apply the law of a foreign nation or to
6 dismiss a claim of a foreign nation or of a citizen
7 of a foreign nation on the ground of inconvenient
8 forum.

9 **SEC. 403. FEDERAL CAUSE OF ACTION PRECLUDED.**

10 The district courts of the United States shall not
11 have jurisdiction pursuant to this Act based on section
12 1331 or 1337 of title 28, United States Code.

13 **SEC. 404. EFFECTIVE DATE.**

14 Titles I, II, and III shall apply with respect to actions
15 which are commenced after the date of the enactment of
16 this Act.

 Passed the House of Representatives March 10,
 1995.

 Attest:

 ROBIN H. CARLE,

Clerk.

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HR 956 PCS—3