

VETO OF H.R. 956

MESSAGE

FROM

THE PRESIDENT OF THE UNITED STATES

TRANSMITTING

HIS VETO OF H.R. 956, A BILL TO ESTABLISH LEGAL STANDARDS
AND PROCEDURES FOR PRODUCT LIABILITY LITIGATION, AND
FOR OTHER PURPOSES



MAY 6, 1996.—Message and accompanying bill ordered to be printed

U.S. GOVERNMENT PRINTING OFFICE

To The House of Representatives:

I am returning herewith without my approval H.R. 956, the "Common Sense Product Liability Legal Reform Act of 1996."

I support real commonsense product liability reform. To deserve that label, however, legislation must adequately protect the interests of consumers, in addition to the interests of manufacturers and sellers. Further, the legislation must respect the important role of the States in our Federal system. The Congress could have passed such legislation, appropriately limited in scope and balanced in application, meeting these tests. Had the Congress done so, I would have signed the bill gladly. The Congress, however, chose not to do so, deciding instead to retain provisions in the bill that I made clear I could not accept.

This bill inappropriately intrudes on State authority, and does so in a way that tilts the legal playing field against consumers. While some Federal action in this area is proper because no one State can alleviate nationwide problems in the tort system, the States should have, as they always have had, primary responsibility for tort law. The States traditionally have handled this job well, serving as laboratories for new ideas and making needed reforms. This bill unduly interferes with that process in products cases; moreover, it does so in a way that peculiarly disadvantages consumers. As a rule, this bill displaces State law only when that law is more favorable to consumers; it defers to State law when that law is more helpful to manufacturers and sellers. I cannot accept, absent compelling reasons, such a one-way street of federalism.

Apart from this general problem of displacing State authority in an unbalanced manner, specific provisions of H.R. 956 unfairly disadvantage consumers and their families. Consumers should be able to count on the safety of the products they purchase. And if these products are defective and cause harm, consumers should be able to get adequate compensation for their losses. Certain provisions in this bill work against these goals, preventing some injured persons from recovering the full measure of their damages and increasing the possibility that defective goods will come onto the market as a result of intentional misconduct.

In particular, I object to the following provisions of the bill, which subject consumers to too great a risk of harm.

First, as I previously have stated, I oppose wholly eliminating joint liability for noneconomic damages such as pain and suffering because such a change would prevent many persons from receiving full compensation for injury. When one wrongdoer cannot pay its portion of the judgment, the other wrongdoers, and not the innocent victim, should have to shoulder that part of the award. Traditional law accomplishes this result. In contrast, this bill would leave the victim to bear these damages on his or her own. Given

how often companies that manufacture defective products go bankrupt, this provision has potentially large consequences.

This provision is all the more troubling because it unfairly discriminates against the most vulnerable members of our society—the elderly, the poor, children, and nonworking women—whose injuries often involve mostly noneconomic losses. There is no reason for this kind of discrimination. Noneconomic damages are as real and as important to victims as economic damages. We should not create a tort system in which people with the greatest need of protection stand the least chance of receiving it.

Second, as I also have stated, I oppose arbitrary ceilings on punitive damages, because they endanger the safety of the public. Capping punitive damages undermines their very purpose, which is to punish and thereby deter egregious misconduct. The provision of the bill allowing judges to exceed the cap if certain factors are present helps to mitigate, but does not cure this problem, given the clear intent of the Congress, as expressed in the Statement of Managers, that judges should use this authority only in the most unusual cases.

In addition, I am concerned that the Conference Report fails to fix an oversight in title II of the bill, which limits actions against suppliers of materials used in devices implanted in the body. In general, title II is a laudable attempt to ensure the supply of materials needed to make life-saving medical devices, such as artificial heart valves. But as I believe even many supporters of the bill agree, a supplier of materials who knew or should have known that the materials, as implanted, would cause injury should not receive any protection from suit. Title II's protections must be clearly limited to nonnegligent suppliers.

My opposition to these Senate-passed provisions were known prior to the Conference on the bill. But instead of addressing these issues, the Conference Committee took several steps backward in the direction of the bill approved by the House.

First, the Conference Report seems to expand the scope of the bill, inappropriately applying the limits on punitive and noneconomic damages to lawsuits, where, for example, a gun dealer has knowingly sold a gun to a convicted felon or a bar owner has knowingly served a drink to an obviously inebriated customer. I believe that such suits should go forward unhindered. Some in the Congress have argued that the change made in Conference is technical in nature, so that the bill still exempts these actions. But I do not read the change in this way—and in any event, I do not believe that a victim of a drunk driver should have to argue in court about this matter. The Congress should not have made this last-minute change, creating this unfortunate ambiguity, in the scope of the bill.

In addition, the Conference Report makes certain changes that, though sounding technical, may cut off a victim's ability to sue a negligent manufacturer. The Report deletes a provision that would have stopped the statute of limitations from running when a bankruptcy court issues the automatic stay that prevents suits from being filed during bankruptcy proceedings. The effect of this seemingly legalistic change will be that some persons harmed by companies that have entered bankruptcy proceedings (as makers of defec-

tive products often do) will lose any meaningful opportunity to bring valid claims.

Similarly, the Conference Report reduces the statute of repose to 15 years (and less if States so provide) and applies the statute to a wider range of goods, including handguns. This change, which bars a suit against a maker of an older product even if that product has just caused injury, also will preclude some valid suits.

In recent weeks, I have heard from many victims of defective products whose efforts to recover compensation would have been frustrated by this bill. I have heard from a woman who would not have received full compensatory damages under this bill for the death of a child because one wrongdoer could not pay his portion of the judgment. I have heard from women whose suits against makers of defective contraceptive devices—and the punitive damages awarded in those suits—forced the products off the market, in a way that this bill's cap on punitives would make much harder. I have heard from persons injured by products more than 15 years old, who under this bill could not bring suit at all.

Injured people cannot be left to suffer in this fashion; furthermore, the few companies that cause these injuries cannot be left, through lack of a deterrent, to engage in misconduct. I therefore must return the bill that has been presented to me. This bill would undermine the ability of courts to provide relief to victims of harmful products and thereby endanger the health and safety of the entire American public. There is nothing common sense about such reforms to product liability law.

WILLIAM J. CLINTON.

THE WHITE HOUSE, *May 2, 1996.*

H.R. 956

ONE HUNDRED FOURTH CONGRESS OF THE UNITED STATES OF AMERICA, AT THE FIRST SESSION, BEGUN AND HELD AT THE CITY OF WASHINGTON ON WEDNESDAY, THE THIRD DAY OF JANUARY, ONE THOUSAND NINE HUNDRED AND NINETY-SIX

An Act

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

Be it enacted by the Senate and House of Representatives of the United States of America in Congress assembled,

SECTION 1. SHORT TITLE AND TABLE OF CONTENTS.

(a) SHORT TITLE.—This Act may be cited as the “Common Sense Product Liability Legal Reform Act of 1996”.

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

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

TITLE I—PRODUCT LIABILITY REFORM

- Sec. 101. Definitions.
- Sec. 102. Applicability; preemption.
- Sec. 103. Liability rules applicable to product sellers, renters, and lessors.
- Sec. 104. Defense based on claimant’s use of intoxicating alcohol or drugs.
- Sec. 105. Misuse or alteration.
- Sec. 106. Uniform time limitations on liability.
- Sec. 107. Alternative dispute resolution procedures.
- Sec. 108. Uniform standards for award of punitive damages.
- Sec. 109. Liability for certain claims relating to death.
- Sec. 110. Several liability for noneconomic loss.
- Sec. 111. Workers’ compensation subrogation.

TITLE II—BIOMATERIALS ACCESS ASSURANCE

- Sec. 201. Short title.
- Sec. 202. Findings.
- Sec. 203. Definitions.
- Sec. 204. General requirements; applicability; preemption.
- Sec. 205. Liability of biomaterials suppliers.
- Sec. 206. Procedures for dismissal of civil actions against biomaterials suppliers.

TITLE III—LIMITATIONS ON APPLICABILITY; EFFECTIVE DATE

- Sec. 301. Effect of court of appeals decisions.
- Sec. 302. Federal cause of action precluded.
- Sec. 303. Effective date.

SEC. 2. FINDINGS AND PURPOSES.

(a) FINDINGS.—The Congress finds that—

(1) our Nation is overly litigious, the civil justice system is overcrowded, sluggish, and excessively costly and the costs of lawsuits, both direct and indirect, are inflicting serious and unnecessary injury on the national economy;

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

(3) the rules of law governing product liability actions, damage awards, and allocations of liability have evolved

inconsistently within and among the States, resulting in a complex, contradictory, and uncertain regime that is inequitable to both plaintiffs and defendants and unduly burdens interstate commerce;

(4) as a result of excessive, unpredictable, and often arbitrary damage awards and unfair allocations of liability, consumers have been adversely affected through the withdrawal of products, producers, services, and service providers from the marketplace, and from excessive liability costs passed on to them through higher prices;

(5) excessive, unpredictable, and often arbitrary damage awards and unfair allocations of liability jeopardize the financial well-being of many individuals as well as entire industries, particularly the Nation's small businesses and adversely affects government and taxpayers;

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

(7) the unpredictability of damage awards is inequitable to both plaintiffs and defendants and has added considerably to the high cost of liability insurance, making it difficult for producers, consumers, volunteers, and nonprofit organizations to protect themselves from liability with any degree of confidence and at a reasonable cost;

(8) because of the national scope of the problems created by the defects in the civil justice system, it is not possible for the States to enact laws that fully and effectively respond to those problems;

(9) it is the constitutional role of the national government to remove barriers to interstate commerce and to protect due process rights; and

(10) there is a need to restore rationality, certainty, and fairness to the civil justice system in order to protect against excessive, arbitrary, and uncertain damage awards and to reduce the volume, costs, and delay of litigation.

(b) PURPOSES.—Based upon the powers contained in Article I, Section 8, Clause 3 and the Fourteenth Amendment of the United States Constitution, the purposes of this Act are to promote the free flow of goods and services and to lessen burdens on interstate commerce and to uphold constitutionally protected due process rights by—

(1) establishing certain uniform legal principles of product liability which provide a fair balance among the interests of product users, manufacturers, and product sellers;

(2) placing reasonable limits on damages over and above the actual damages suffered by a claimant;

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

(4) reducing the unacceptable costs and delays of our civil justice system caused by excessive litigation which harm both plaintiffs and defendants; and

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

TITLE I—PRODUCT LIABILITY REFORM

SEC. 101. DEFINITIONS.

For purposes of this title—

(1) **ACTUAL MALICE.**—The term “actual malice” means specific intent to cause serious physical injury, illness, disease, death, or damage to property.

(2) **CLAIMANT.**—The term “claimant” means any person who brings an action covered by this title and any person on whose behalf such an action is brought. If such an action is brought through or on behalf of an estate, the term includes the claimant’s decedent. If such an action is brought through or on behalf of a minor or incompetent, the term includes the claimant’s legal guardian.

(3) **CLAIMANT’S BENEFITS.**—The term “claimant’s benefits” means the amount paid to an employee as workers’ compensation benefits.

(4) **CLEAR AND CONVINCING EVIDENCE.**—The term “clear and convincing evidence” is that measure or degree of proof that will produce in the mind of the trier of fact a firm belief or conviction as to the truth of the allegations sought to be established. The level of proof required to satisfy such standard is more than that required under preponderance of the evidence, but less than that required for proof beyond a reasonable doubt.

(5) **COMMERCIAL LOSS.**—The term “commercial loss” means any loss or damage solely to a product itself, loss relating to a dispute over its value, or consequential economic loss, the recovery of which is governed by the Uniform Commercial Code or analogous State commercial or contract law.

(6) **COMPENSATORY DAMAGES.**—The term “compensatory damages” means damages awarded for economic and non-economic loss.

(7) **DURABLE GOOD.**—The term “durable good” means any product, or any component of any such product, which has a normal life expectancy of 3 or more years, or is of a character subject to allowance for depreciation under the Internal Revenue Code of 1986 and which is—

(A) used in a trade or business;

(B) held for the production of income; or

(C) sold or donated to a governmental or private entity for the production of goods, training, demonstration, or any other similar purpose.

(8) **ECONOMIC LOSS.**—The term “economic loss” means any pecuniary loss resulting from harm (including the loss of earnings or other benefits related to employment, medical expense loss, replacement services loss, loss due to death, burial costs, and loss of business or employment opportunities) to the extent recovery for such loss is allowed under applicable State law.

(9) **HARM.**—The term “harm” means any physical injury, illness, disease, or death or damage to property caused by a product. The term does not include commercial loss.

(10) **INSURER.**—The term “insurer” means the employer of a claimant if the employer is self-insured or if the employer is not self-insured, the workers’ compensation insurer of the employer.

(11) **MANUFACTURER.**—The term “manufacturer” means—

(A) any person who is engaged in a business to produce, create, make, or construct any product (or component part of a product) and who (i) designs or formulates the product (or component part of the product), or (ii) has engaged another person to design or formulate the product (or component part of the product);

(B) a product seller, but only with respect to those aspects of a product (or component part of a product) which are created or affected when, before placing the product in the stream of commerce, the product seller produces, creates, makes or constructs and designs, or formulates, or has engaged another person to design or formulate, an aspect of the product (or component part of the product) made by another person; or

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

(12) NONECONOMIC LOSS.—The term “noneconomic loss” means subjective, nonmonetary loss resulting from harm, including pain, suffering, inconvenience, mental suffering, emotional distress, loss of society and companionship, loss of consortium, injury to reputation, and humiliation.

(13) PERSON.—The term “person” means any individual, corporation, company, association, firm, partnership, society, joint stock company, or any other entity (including any governmental entity).

(14) PRODUCT.—

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

(i) is capable of delivery itself or as an assembled whole, in a mixed or combined state, or as a component part or ingredient;

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

(iii) has intrinsic economic value; and

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

(B) EXCLUSION.—The term does not include—

(i) tissue, organs, blood, and blood products used for therapeutic or medical purposes, except to the extent that such tissue, organs, blood, and blood products (or the provision thereof) are subject, under applicable State law, to a standard of liability other than negligence; or

(ii) electricity, water delivered by a utility, natural gas, or steam except to the extent that electricity, water delivered by a utility, natural gas, or steam, is subject, under applicable State law, to a standard of liability other than negligence.

(15) PRODUCT LIABILITY ACTION.—The term “product liability action” means a civil action brought on any theory for harm caused by a product.

(16) PRODUCT SELLER.—

(A) IN GENERAL.—The term “product seller” means a person who in the course of a business conducted for that purpose—

(i) sells, distributes, rents, leases, prepares, blends, packages, labels, or otherwise is involved in placing a product in the stream of commerce; or

(ii) installs, repairs, refurbishes, reconditions, or maintains the harm-causing aspect of the product.

(B) EXCLUSION.—The term “product seller” does not include—

(i) a seller or lessor of real property;

(ii) a provider of professional services in any case in which the sale or use of a product is incidental to the transaction and the essence of the transaction is the furnishing of judgment, skill, or services; or

(iii) any person who—

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

(II) leases a product under a lease arrangement in which the lessor does not initially select the leased product and does not during the lease term ordinarily control the daily operations and maintenance of the product.

(17) PUNITIVE DAMAGES.—The term “punitive damages” means damages awarded against any person or entity to punish or deter such person or entity, or others, from engaging in similar behavior in the future.

(18) STATE.—The term “State” means any State of the United States, the District of Columbia, Commonwealth of Puerto Rico, the Northern Mariana Islands, the Virgin Islands, Guam, American Samoa, and any other territory or possession of the United States or any political subdivision of any of the foregoing.

SEC. 102. APPLICABILITY; PREEMPTION.

(a) PREEMPTION.—

(1) IN GENERAL.—This Act governs any product liability action brought in any State or Federal court on any theory for harm caused by a product.

(2) ACTIONS EXCLUDED.—A civil action brought for commercial loss shall be governed only by applicable commercial or contract law.

(b) RELATIONSHIP TO STATE LAW.—This title supersedes State law only to the extent that State law applies to an issue covered by this title. Any issue that is not governed by this title, including any standard of liability applicable to a manufacturer, shall be governed by otherwise applicable State or Federal law.

(c) EFFECT ON OTHER LAW.—Nothing in this Act shall be construed to—

(1) waive or affect any defense of sovereign immunity asserted by any State under any law;

(2) supersede or alter any Federal law;

(3) waive or affect any defense of sovereign immunity asserted by the United States;

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

(5) preempt State choice-of-law rules with respect to claims brought by a foreign nation or a citizen of a foreign nation;

(6) affect the right of any court to transfer venue or to apply the law of a foreign nation or to dismiss a claim of

a foreign nation or of a citizen of a foreign nation on the ground of inconvenient forum; or

(7) supersede or modify any statutory or common law, including any law providing for an action to abate a nuisance, that authorizes a person to institute an action for civil damages or civil penalties, cleanup costs, injunctions, restitution, cost recovery, punitive damages, or any other form of relief for remediation of the environment (as defined in section 101(8) of the Comprehensive Environmental Response, Compensation, and Liability Act of 1980 (42 U.S.C. 9601(8)).

SEC. 103. LIABILITY RULES APPLICABLE TO PRODUCT SELLERS, RENTERS, AND LESSORS.

(a) GENERAL RULE.—

(1) IN GENERAL.—In any product liability action, a product seller other than a manufacturer shall be liable to a claimant only if the claimant establishes—

(A) that—

(i) the product that allegedly caused the harm that is the subject of the complaint was sold, rented, or leased by the product seller;

(ii) the product seller failed to exercise reasonable care with respect to the product; and

(iii) the failure to exercise reasonable care was a proximate cause of harm to the claimant;

(B) that—

(i) the product seller made an express warranty applicable to the product that allegedly caused the harm that is the subject of the complaint, independent of any express warranty made by a manufacturer as to the same product;

(ii) the product failed to conform to the warranty; and

(iii) the failure of the product to conform to the warranty caused harm to the claimant; or

(C) that—

(i) the product seller engaged in intentional wrongdoing, as determined under applicable State law; and

(ii) such intentional wrongdoing was a proximate cause of the harm that is the subject of the complaint.

(2) REASONABLE OPPORTUNITY FOR INSPECTION.—For purposes of paragraph (1)(A)(ii), a product seller shall not be considered to have failed to exercise reasonable care with respect to a product based upon an alleged failure to inspect the product—

(A) if the failure occurred because there was no reasonable opportunity to inspect the product; or

(B) if the inspection, in the exercise of reasonable care, would not have revealed the aspect of the product which allegedly caused the claimant's harm.

(b) SPECIAL RULE.—

(1) IN GENERAL.—A product seller shall be deemed to be liable as a manufacturer of a product for harm caused by the product if—

(A) the manufacturer is not subject to service of process under the laws of any State in which the action may be brought; or

(B) the court determines that the claimant would be unable to enforce a judgment against the manufacturer.

(2) **STATUTE OF LIMITATIONS.**—For purposes of this subsection only, the statute of limitations applicable to claims asserting liability of a product seller as a manufacturer shall be tolled from the date of the filing of a complaint against the manufacturer to the date that judgment is entered against the manufacturer.

(c) **RENTED OR LEASED PRODUCTS.**—

(1) Notwithstanding any other provision of law, any person engaged in the business of renting or leasing a product (other than a person excluded from the definition of product seller under section 101(16)(B)) shall be subject to liability in a product liability action under subsection (a), but any person engaged in the business of renting or leasing a product shall not be liable to a claimant for the tortious act of another solely by reason of ownership of such product.

(2) For purposes of paragraph (1), and for determining the applicability of this title to any person subject to paragraph (1), the term “product liability action” means a civil action brought on any theory for harm caused by a product or product use.

(d) **ACTIONS FOR NEGLIGENT ENTRUSTMENT.**—A civil action for negligent entrustment shall not be subject to the provisions of this section, but shall be subject to any applicable State law.

SEC. 104. DEFENSE BASED ON CLAIMANT’S USE OF INTOXICATING ALCOHOL OR DRUGS.

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

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

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

(b) **CONSTRUCTION.**—For purposes of subsection (a)—

(1) the determination of whether a person was intoxicated or was under the influence of intoxicating alcohol or any drug shall be made pursuant to applicable State law; and

(2) the term “drug” means any controlled substance as defined in the Controlled Substances Act (21 U.S.C. 802(6)) that was not legally prescribed for use by the claimant or that was taken by the claimant other than in accordance with the terms of a lawfully issued prescription.

SEC. 105. MISUSE OR ALTERATION.

(a) **GENERAL RULE.**—

(1) **IN GENERAL.**—In a product liability action, the damages for which a defendant is otherwise liable under Federal or State law shall be reduced by the percentage of responsibility for the claimant’s harm attributable to misuse or alteration of a product by any person if the defendant establishes that such percentage of the claimant’s harm was proximately caused by a use or alteration of a product—

(A) in violation of, or contrary to, a defendant’s express warnings or instructions if the warnings or instructions

are adequate as determined pursuant to applicable State law; or

(B) involving a risk of harm which was known or should have been known by the ordinary person who uses or consumes the product with the knowledge common to the class of persons who used or would be reasonably anticipated to use the product.

(2) USE INTENDED BY A MANUFACTURER IS NOT MISUSE OR ALTERATION.—For the purposes of this Act, a use of a product that is intended by the manufacturer of the product does not constitute a misuse or alteration of the product.

(b) WORKPLACE INJURY.—Notwithstanding subsection (a), and except as otherwise provided in section 111, the damages for which a defendant is otherwise liable under State law shall not be reduced by the percentage of responsibility for the claimant's harm attributable to misuse or alteration of the product by the claimant's employer or any coemployee who is immune from suit by the claimant pursuant to the State law applicable to workplace injuries.

SEC. 106. UNIFORM TIME LIMITATIONS ON LIABILITY.

(a) STATUTE OF LIMITATIONS.—

(1) IN GENERAL.—Except as provided in paragraph (2) and subsection (b), a product liability action may be filed not later than 2 years after the date on which the claimant discovered or, in the exercise of reasonable care, should have discovered—

- (A) the harm that is the subject of the action; and
- (B) the cause of the harm.

(2) EXCEPTION.—A person with a legal disability (as determined under applicable law) may file a product liability action not later than 2 years after the date on which the person ceases to have the legal disability.

(b) STATUTE OF REPOSE.—

(1) IN GENERAL.—Subject to paragraphs (2) and (3), no product liability action that is subject to this Act concerning a product, that is a durable good, alleged to have caused harm (other than toxic harm) may be filed after the 15-year period beginning at the time of delivery of the product to the first purchaser or lessee.

(2) STATE LAW.—Notwithstanding paragraph (1), if pursuant to an applicable State law, an action described in such paragraph is required to be filed during a period that is shorter than the 15-year period specified in such paragraph, the State law shall apply with respect to such period.

(3) EXCEPTIONS.—

(A) A motor vehicle, vessel, aircraft, or train, that is used primarily to transport passengers for hire, shall not be subject to this subsection.

(B) Paragraph (1) does not bar a product liability action against a defendant who made an express warranty in writing as to the safety or life expectancy of the specific product involved which was longer than 15 years, but it will apply at the expiration of that warranty.

(C) Paragraph (1) does not affect the limitations period established by the General Aviation Revitalization Act of 1994 (49 U.S.C. 40101 note).

(c) TRANSITIONAL PROVISION RELATING TO EXTENSION OF PERIOD FOR BRINGING CERTAIN ACTIONS.—If any provision of sub-

section (a) or (b) shortens the period during which a product liability action could be otherwise brought pursuant to another provision of law, the claimant may, notwithstanding subsections (a) and (b), bring the product liability action not later than 1 year after the date of enactment of this Act.

SEC. 107. ALTERNATIVE DISPUTE RESOLUTION PROCEDURES.

(a) SERVICE OF OFFER.—A claimant or a defendant in a product liability action may, not later than 60 days after the service of—

- (1) the initial complaint; or
- (2) the applicable deadline for a responsive pleading;

whichever is later, serve upon an adverse party an offer to proceed pursuant to any voluntary, nonbinding alternative dispute resolution procedure established or recognized under the law of the State in which the product liability action is brought or under the rules of the court in which such action is maintained.

(b) WRITTEN NOTICE OF ACCEPTANCE OR REJECTION.—Except as provided in subsection (c), not later than 10 days after the service of an offer to proceed under subsection (a), an offeree shall file a written notice of acceptance or rejection of the offer.

(c) EXTENSION.—The court may, upon motion by an offeree made prior to the expiration of the 10-day period specified in subsection (b), extend the period for filing a written notice under such subsection for a period of not more than 60 days after the date of expiration of the period specified in subsection (b). Discovery may be permitted during such period.

SEC. 108. UNIFORM STANDARDS FOR AWARD OF PUNITIVE DAMAGES.

(a) GENERAL RULE.—Punitive damages may, to the extent permitted by applicable State law, be awarded against a defendant if the claimant establishes by clear and convincing evidence that conduct carried out by the defendant with a conscious, flagrant indifference to the rights or safety of others was the proximate cause of the harm that is the subject of the action in any product liability action.

(b) LIMITATION ON AMOUNT.—

(1) IN GENERAL.—The amount of punitive damages that may be awarded in an action described in subsection (a) may not exceed the greater of—

- (A) 2 times the sum of the amount awarded to the claimant for economic loss and noneconomic loss; or
- (B) \$250,000.

(2) SPECIAL RULE.—Notwithstanding paragraph (1), in any action described in subsection (a) against an individual whose net worth does not exceed \$500,000 or against an owner of an unincorporated business, or any partnership, corporation, association, unit of local government, or organization which has fewer than 25 full-time employees, the punitive damages shall not exceed the lesser of—

- (A) 2 times the sum of the amount awarded to the claimant for economic loss and noneconomic loss; or
- (B) \$250,000.

For the purpose of determining the applicability of this paragraph to a corporation, the number of employees of a subsidiary or wholly-owned corporation shall include all employees of a parent or sister corporation.

(3) EXCEPTION FOR INSUFFICIENT AWARD IN CASES OF EGREGIOUS CONDUCT.—

(A) DETERMINATION BY COURT.—If the court makes a determination, after considering each of the factors in subparagraph (B), that the application of paragraph (1) would result in an award of punitive damages that is insufficient to punish the egregious conduct of the defendant against whom the punitive damages are to be awarded or to deter such conduct in the future, the court shall determine the additional amount of punitive damages (referred to in this paragraph as the “additional amount”) in excess of the amount determined in accordance with paragraph (1) to be awarded against the defendant in a separate proceeding in accordance with this paragraph.

(B) FACTORS FOR CONSIDERATION.—In any proceeding under paragraph (A), the court shall consider—

(i) the extent to which the defendant acted with actual malice;

(ii) the likelihood that serious harm would arise from the conduct of the defendant;

(iii) the degree of the awareness of the defendant of that likelihood;

(iv) the profitability of the misconduct to the defendant;

(v) the duration of the misconduct and any concurrent or subsequent concealment of the conduct by the defendant;

(vi) the attitude and conduct of the defendant upon the discovery of the misconduct and whether the misconduct has terminated;

(vii) the financial condition of the defendant; and

(viii) the cumulative deterrent effect of other losses, damages, and punishment suffered by the defendant as a result of the misconduct, reducing the amount of punitive damages on the basis of the economic impact and severity of all measures to which the defendant has been or may be subjected, including—

(I) compensatory and punitive damage awards to similarly situated claimants;

(II) the adverse economic effect of stigma or loss of reputation;

(III) civil fines and criminal and administrative penalties; and

(IV) stop sale, cease and desist, and other remedial or enforcement orders.

(C) REQUIREMENTS FOR AWARDED ADDITIONAL AMOUNT.—If the court awards an additional amount pursuant to this subsection, the court shall state its reasons for setting the amount of the additional amount in findings of fact and conclusions of law.

(D) PREEMPTION.—This section does not create a cause of action for punitive damages and does not preempt or supersede any State or Federal law to the extent that such law would further limit the award of punitive damages. Nothing in this subsection shall modify or reduce the ability of courts to order remittiturs.

(4) APPLICATION BY COURT.—This subsection shall be applied by the court and application of this subsection shall not be disclosed to the jury. Nothing in this subsection shall authorize the court to enter an award of punitive damages in excess of the jury's initial award of punitive damages.

(c) BIFURCATION AT REQUEST OF ANY PARTY.—

(1) IN GENERAL.—At the request of any party the trier of fact in any action that is subject to this section shall consider in a separate proceeding, held subsequent to the determination of the amount of compensatory damages, whether punitive damages are to be awarded for the harm that is the subject of the action and the amount of the award.

(2) INADMISSIBILITY OF EVIDENCE RELATIVE ONLY TO A CLAIM OF PUNITIVE DAMAGES IN A PROCEEDING CONCERNING COMPENSATORY DAMAGES.—If any party requests a separate proceeding under paragraph (1), in a proceeding to determine whether the claimant may be awarded compensatory damages, any evidence, argument, or contention that is relevant only to the claim of punitive damages, as determined by applicable State law, shall be inadmissible.

SEC. 109. LIABILITY FOR CERTAIN CLAIMS RELATING TO DEATH.

In any civil action in which the alleged harm to the claimant is death and, as of the effective date of this Act, the applicable State law provides, or has been construed to provide, for damages only punitive in nature, a defendant may be liable for any such damages without regard to section 108, but only during such time as the State law so provides. This section shall cease to be effective September 1, 1996.

SEC. 110. SEVERAL LIABILITY FOR NONECONOMIC LOSS.

(a) GENERAL RULE.—In a product liability action, the liability of each defendant for noneconomic loss shall be several only and shall not be joint.

(b) AMOUNT OF LIABILITY.—

(1) IN GENERAL.—Each defendant shall be liable only for the amount of noneconomic loss allocated to the defendant in direct proportion to the percentage of responsibility of the defendant (determined in accordance with paragraph (2)) for the harm to the claimant with respect to which the defendant is liable. The court shall render a separate judgment against each defendant in an amount determined pursuant to the preceding sentence.

(2) PERCENTAGE OF RESPONSIBILITY.—For purposes of determining the amount of noneconomic loss allocated to a defendant under this section, the trier of fact shall determine the percentage of responsibility of each person responsible for the claimant's harm, whether or not such person is a party to the action.

SEC. 111. WORKERS' COMPENSATION SUBROGATION.

(a) GENERAL RULE.—

(1) RIGHT OF SUBROGATION.—

(A) IN GENERAL.—An insurer shall have a right of subrogation against a manufacturer or product seller to recover any claimant's benefits relating to harm that is the subject of a product liability action that is subject to this Act.

(B) WRITTEN NOTIFICATION.—To assert a right of subrogation under subparagraph (A), the insurer shall provide written notice to the court in which the product liability action is brought.

(C) INSURER NOT REQUIRED TO BE A PARTY.—An insurer shall not be required to be a necessary and proper party in a product liability action covered under subparagraph (A).

(2) SETTLEMENTS AND OTHER LEGAL PROCEEDINGS.—

(A) IN GENERAL.—In any proceeding relating to harm or settlement with the manufacturer or product seller by a claimant who files a product liability action that is subject to this Act, an insurer may participate to assert a right of subrogation for claimant's benefits with respect to any payment made by the manufacturer or product seller by reason of such harm, without regard to whether the payment is made—

- (i) as part of a settlement;
- (ii) in satisfaction of judgment;
- (iii) as consideration for a covenant not to sue;

or

- (iv) in another manner.

(B) WRITTEN NOTIFICATION.—Except as provided in subparagraph (C), an employee shall not make any settlement with or accept any payment from the manufacturer or product seller without written notification to the insurer.

(C) EXEMPTION.—Subparagraph (B) shall not apply in any case in which the insurer has been compensated for the full amount of the claimant's benefits.

(3) HARM RESULTING FROM ACTION OF EMPLOYER OR CO-EMPLOYEE.—

(A) IN GENERAL.—If, with respect to a product liability action that is subject to this Act, the manufacturer or product seller attempts to persuade the trier of fact that the harm to the claimant was caused by the fault of the employer of the claimant or any coemployee of the claimant, the issue of that fault shall be submitted to the trier of fact, but only after the manufacturer or product seller has provided timely written notice to the insurer.

(B) RIGHTS OF INSURER.—

(i) IN GENERAL.—Notwithstanding any other provision of law, with respect to an issue of fault submitted to a trier of fact pursuant to subparagraph (A), an insurer shall, in the same manner as any party in the action (even if the insurer is not a named party in the action), have the right to—

- (I) appear;
- (II) be represented;
- (III) introduce evidence;
- (IV) cross-examine adverse witnesses; and
- (V) present arguments to the trier of fact.

(ii) LAST ISSUE.—The issue of harm resulting from an action of an employer or coemployee shall be the last issue that is submitted to the trier of fact.

(C) REDUCTION OF DAMAGES.—If the trier of fact finds by clear and convincing evidence that the harm to the claimant that is the subject of the product liability action

was caused by the fault of the employer or a coemployee of the claimant—

(i) the court shall reduce by the amount of the claimant's benefits—

(I) the damages awarded against the manufacturer or product seller; and

(II) any corresponding insurer's subrogation lien; and

(ii) the manufacturer or product seller shall have no further right by way of contribution or otherwise against the employer.

(D) CERTAIN RIGHTS OF SUBROGATION NOT AFFECTED.—Notwithstanding a finding by the trier of fact described in subparagraph (C), the insurer shall not lose any right of subrogation related to any—

(i) intentional tort committed against the claimant by a coemployee; or

(ii) act committed by a coemployee outside the scope of normal work practices.

(b) ATTORNEY'S FEES.—If, in a product liability action that is subject to this section, the court finds that harm to a claimant was not caused by the fault of the employer or a coemployee of the claimant, the manufacturer or product seller shall reimburse the insurer for reasonable attorney's fees and court costs incurred by the insurer in the action, as determined by the court.

TITLE II—BIOMATERIALS ACCESS ASSURANCE

SEC. 201. SHORT TITLE.

This title may be cited as the "Biomaterials Access Assurance Act of 1996".

SEC. 202. FINDINGS.

Congress finds that—

(1) each year millions of citizens of the United States depend on the availability of lifesaving or life enhancing medical devices, many of which are permanently implantable within the human body;

(2) a continued supply of raw materials and component parts is necessary for the invention, development, improvement, and maintenance of the supply of the devices;

(3) most of the medical devices are made with raw materials and component parts that—

(A) are not designed or manufactured specifically for use in medical devices; and

(B) come in contact with internal human tissue;

(4) the raw materials and component parts also are used in a variety of nonmedical products;

(5) because small quantities of the raw materials and component parts are used for medical devices, sales of raw materials and component parts for medical devices constitute an extremely small portion of the overall market for the raw materials and medical devices;

(6) under the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 301 et seq.), manufacturers of medical devices are

required to demonstrate that the medical devices are safe and effective, including demonstrating that the products are properly designed and have adequate warnings or instructions;

(7) notwithstanding the fact that raw materials and component parts suppliers do not design, produce, or test a final medical device, the suppliers have been the subject of actions alleging inadequate—

(A) design and testing of medical devices manufactured with materials or parts supplied by the suppliers; or

(B) warnings related to the use of such medical devices;

(8) even though suppliers of raw materials and component parts have very rarely been held liable in such actions, such suppliers have ceased supplying certain raw materials and component parts for use in medical devices because the costs associated with litigation in order to ensure a favorable judgment for the suppliers far exceeds the total potential sales revenues from sales by such suppliers to the medical device industry;

(9) unless alternate sources of supply can be found, the unavailability of raw materials and component parts for medical devices will lead to unavailability of lifesaving and life-enhancing medical devices;

(10) because other suppliers of the raw materials and component parts in foreign nations are refusing to sell raw materials or component parts for use in manufacturing certain medical devices in the United States, the prospects for development of new sources of supply for the full range of threatened raw materials and component parts for medical devices are remote;

(11) it is unlikely that the small market for such raw materials and component parts in the United States could support the large investment needed to develop new suppliers of such raw materials and component parts;

(12) attempts to develop such new suppliers would raise the cost of medical devices;

(13) courts that have considered the duties of the suppliers of the raw materials and component parts have generally found that the suppliers do not have a duty—

(A) to evaluate the safety and efficacy of the use of a raw material or component part in a medical device; and

(B) to warn consumers concerning the safety and effectiveness of a medical device;

(14) attempts to impose the duties referred to in subparagraphs (A) and (B) of paragraph (13) on suppliers of the raw materials and component parts would cause more harm than good by driving the suppliers to cease supplying manufacturers of medical devices; and

(15) in order to safeguard the availability of a wide variety of lifesaving and life-enhancing medical devices, immediate action is needed—

(A) to clarify the permissible bases of liability for suppliers of raw materials and component parts for medical devices; and

(B) to provide expeditious procedures to dispose of unwarranted suits against the suppliers in such manner as to minimize litigation costs.

SEC. 203. DEFINITIONS.

As used in this title:

(1) BIOMATERIALS SUPPLIER.—

(A) **IN GENERAL.**—The term “biomaterials supplier” means an entity that directly or indirectly supplies a component part or raw material for use in the manufacture of an implant.

(B) **PERSONS INCLUDED.**—Such term includes any person who—

(i) has submitted master files to the Secretary for purposes of premarket approval of a medical device; or

(ii) licenses a biomaterials supplier to produce component parts or raw materials.

(2) CLAIMANT.—

(A) **IN GENERAL.**—The term “claimant” means any person who brings a civil action, or on whose behalf a civil action is brought, arising from harm allegedly caused directly or indirectly by an implant, including a person other than the individual into whose body, or in contact with whose blood or tissue, the implant is placed, who claims to have suffered harm as a result of the implant.

(B) **ACTION BROUGHT ON BEHALF OF AN ESTATE.**—With respect to an action brought on behalf of or through the estate of an individual into whose body, or in contact with whose blood or tissue the implant is placed, such term includes the decedent that is the subject of the action.

(C) **ACTION BROUGHT ON BEHALF OF A MINOR OR INCOMPETENT.**—With respect to an action brought on behalf of or through a minor or incompetent, such term includes the parent or guardian of the minor or incompetent.

(D) **EXCLUSIONS.**—Such term does not include—

(i) a provider of professional health care services, in any case in which—

(I) the sale or use of an implant is incidental to the transaction; and

(II) the essence of the transaction is the furnishing of judgment, skill, or services; or

(ii) a person acting in the capacity of a manufacturer, seller, or biomaterials supplier.

(3) COMPONENT PART.—

(A) **IN GENERAL.**—The term “component part” means a manufactured piece of an implant.

(B) **CERTAIN COMPONENTS.**—Such term includes a manufactured piece of an implant that—

(i) has significant non-implant applications; and

(ii) alone, has no implant value or purpose, but when combined with other component parts and materials, constitutes an implant.

(4) HARM.—

(A) **IN GENERAL.**—The term “harm” means—

(i) any injury to or damage suffered by an individual;

(ii) any illness, disease, or death of that individual resulting from that injury or damage; and

(iii) any loss to that individual or any other individual resulting from that injury or damage.

(B) EXCLUSION.—The term does not include any commercial loss or loss of or damage to an implant.

(5) IMPLANT.—The term “implant” means—

(A) a medical device that is intended by the manufacturer of the device—

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

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

(B) suture materials used in implant procedures.

(6) MANUFACTURER.—The term “manufacturer” means any person who, with respect to an implant—

(A) is engaged in the manufacture, preparation, propagation, compounding, or processing (as defined in section 510(a)(1)) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 360(a)(1)) of the implant; and

(B) is required—

(i) to register with the Secretary pursuant to section 510 of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 360) and the regulations issued under such section; and

(ii) to include the implant on a list of devices filed with the Secretary pursuant to section 510(j) of such Act (21 U.S.C. 360(j)) and the regulations issued under such section.

(7) MEDICAL DEVICE.—The term “medical device” means a device, as defined in section 201(h) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 321(h)) and includes any device component of any combination product as that term is used in section 503(g) of such Act (21 U.S.C. 353(g)).

(8) RAW MATERIAL.—The term “raw material” means a substance or product that—

(A) has a generic use; and

(B) may be used in an application other than an implant.

(9) SECRETARY.—The term “Secretary” means the Secretary of Health and Human Services.

(10) SELLER.—

(A) IN GENERAL.—The term “seller” means a person who, in the course of a business conducted for that purpose, sells, distributes, leases, packages, labels, or otherwise places an implant in the stream of commerce.

(B) EXCLUSIONS.—The term does not include—

(i) a seller or lessor of real property;

(ii) a provider of professional services, in any case in which the sale or use of an implant is incidental to the transaction and the essence of the transaction is the furnishing of judgment, skill, or services; or

(iii) any person who acts in only a financial capacity with respect to the sale of an implant.

SEC. 204. GENERAL REQUIREMENTS; APPLICABILITY; PREEMPTION.

(a) GENERAL REQUIREMENTS.—

(1) **IN GENERAL.**—In any civil action covered by this title, a biomaterials supplier may raise any defense set forth in section 205.

(2) **PROCEDURES.**—Notwithstanding any other provision of law, the Federal or State court in which a civil action covered by this title is pending shall, in connection with a motion for dismissal or judgment based on a defense described in paragraph (1), use the procedures set forth in section 206.

(b) **APPLICABILITY.**—

(1) **IN GENERAL.**—Except as provided in paragraph (2), notwithstanding any other provision of law, this title applies to any civil action brought by a claimant, whether in a Federal or State court, against a manufacturer, seller, or biomaterials supplier, on the basis of any legal theory, for harm allegedly caused by an implant.

(2) **EXCLUSION.**—A civil action brought by a purchaser of a medical device for use in providing professional services against a manufacturer, seller, or biomaterials supplier for loss or damage to an implant or for commercial loss to the purchaser—

(A) shall not be considered an action that is subject to this title; and

(B) shall be governed by applicable commercial or contract law.

(c) **SCOPE OF PREEMPTION.**—

(1) **IN GENERAL.**—This title supersedes any State law regarding recovery for harm caused by an implant and any rule of procedure applicable to a civil action to recover damages for such harm only to the extent that this title establishes a rule of law applicable to the recovery of such damages.

(2) **APPLICABILITY OF OTHER LAWS.**—Any issue that arises under this title and that is not governed by a rule of law applicable to the recovery of damages described in paragraph (1) shall be governed by applicable Federal or State law.

(d) **STATUTORY CONSTRUCTION.**—Nothing in this title may be construed—

(1) to affect any defense available to a defendant under any other provisions of Federal or State law in an action alleging harm caused by an implant; or

(2) to create a cause of action or Federal court jurisdiction pursuant to section 1331 or 1337 of title 28, United States Code, that otherwise would not exist under applicable Federal or State law.

SEC. 205. LIABILITY OF BIOMATERIALS SUPPLIERS.

(a) **IN GENERAL.**—

(1) **EXCLUSION FROM LIABILITY.**—Except as provided in paragraph (2), a biomaterials supplier shall not be liable for harm to a claimant caused by an implant.

(2) **LIABILITY.**—A biomaterials supplier that—

(A) is a manufacturer may be liable for harm to a claimant described in subsection (b);

(B) is a seller may be liable for harm to a claimant described in subsection (c); and

(C) furnishes raw materials or component parts that fail to meet applicable contractual requirements or speci-

fications may be liable for a harm to a claimant described in subsection (d).

(b) LIABILITY AS MANUFACTURER.—

(1) IN GENERAL.—A biomaterials supplier may, to the extent required and permitted by any other applicable law, be liable for harm to a claimant caused by an implant if the biomaterials supplier is the manufacturer of the implant.

(2) GROUNDS FOR LIABILITY.—The biomaterials supplier may be considered the manufacturer of the implant that allegedly caused harm to a claimant only if the biomaterials supplier—

(A)(i) has registered with the Secretary pursuant to section 510 of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 360) and the regulations issued under such section; and

(ii) included the implant on a list of devices filed with the Secretary pursuant to section 510(j) of such Act (21 U.S.C. 360(j)) and the regulations issued under such section;

(B) is the subject of a declaration issued by the Secretary pursuant to paragraph (3) that states that the supplier, with respect to the implant that allegedly caused harm to the claimant, was required to—

(i) register with the Secretary under section 510 of such Act (21 U.S.C. 360), and the regulations issued under such section, but failed to do so; or

(ii) include the implant on a list of devices filed with the Secretary pursuant to section 510(j) of such Act (21 U.S.C. 360(j)) and the regulations issued under such section, but failed to do so; or

(C) is related by common ownership or control to a person meeting all the requirements described in subparagraph (A) or (B), if the court deciding a motion to dismiss in accordance with section 206(c)(3)(B)(i) finds, on the basis of affidavits submitted in accordance with section 206, that it is necessary to impose liability on the biomaterials supplier as a manufacturer because the related manufacturer meeting the requirements of subparagraph (A) or (B) lacks sufficient financial resources to satisfy any judgment that the court feels it is likely to enter should the claimant prevail.

(3) ADMINISTRATIVE PROCEDURES.—

(A) IN GENERAL.—The Secretary may issue a declaration described in paragraph (2)(B) on the motion of the Secretary or on petition by any person, after providing—

(i) notice to the affected persons; and

(ii) an opportunity for an informal hearing.

(B) DOCKETING AND FINAL DECISION.—Immediately upon receipt of a petition filed pursuant to this paragraph, the Secretary shall docket the petition. Not later than 180 days after the petition is filed, the Secretary shall issue a final decision on the petition.

(C) APPLICABILITY OF STATUTE OF LIMITATIONS.—Any applicable statute of limitations shall toll during the period during which a claimant has filed a petition with the Secretary under this paragraph.

(c) **LIABILITY AS SELLER.**—A biomaterials supplier may, to the extent required and permitted by any other applicable law, be liable as a seller for harm to a claimant caused by an implant if—

(1) the biomaterials supplier—

(A) held title to the implant that allegedly caused harm to the claimant as a result of purchasing the implant after—

(i) the manufacture of the implant; and

(ii) the entrance of the implant in the stream of commerce; and

(B) subsequently resold the implant; or

(2) the biomaterials supplier is related by common ownership or control to a person meeting all the requirements described in paragraph (1), if a court deciding a motion to dismiss in accordance with section 206(c)(3)(B)(ii) finds, on the basis of affidavits submitted in accordance with section 206, that it is necessary to impose liability on the biomaterials supplier as a seller because the related seller meeting the requirements of paragraph (1) lacks sufficient financial resources to satisfy any judgment that the court feels it is likely to enter should the claimant prevail.

(d) **LIABILITY FOR VIOLATING CONTRACTUAL REQUIREMENTS OR SPECIFICATIONS.**—A biomaterials supplier may, to the extent required and permitted by any other applicable law, be liable for harm to a claimant caused by an implant, if the claimant in an action shows, by a preponderance of the evidence, that—

(1) the raw materials or component parts delivered by the biomaterials supplier either—

(A) did not constitute the product described in the contract between the biomaterials supplier and the person who contracted for delivery of the product; or

(B) failed to meet any specifications that were—

(i) provided to the biomaterials supplier and not expressly repudiated by the biomaterials supplier prior to acceptance of delivery of the raw materials or component parts;

(ii)(I) published by the biomaterials supplier;

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

(III) contained in a master file that was submitted by the biomaterials supplier to the Secretary and that is currently maintained by the biomaterials supplier for purposes of premarket approval of medical devices; or

(iii) included in the submissions for purposes of premarket approval or review by the Secretary under section 510, 513, 515, or 520 of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 360, 360c, 360e, or 360j), and received clearance from the Secretary if such specifications were provided by the manufacturer to the biomaterials supplier and were not expressly repudiated by the biomaterials supplier prior to the acceptance by the manufacturer of delivery of the raw materials or component parts; and

(2) such conduct was an actual and proximate cause of the harm to the claimant.

SEC. 206. PROCEDURES FOR DISMISSAL OF CIVIL ACTIONS AGAINST BIOMATERIALS SUPPLIERS.

(a) **MOTION TO DISMISS.**—In any action that is subject to this title, a biomaterials supplier who is a defendant in such action may, at any time during which a motion to dismiss may be filed under an applicable law, move to dismiss the action against it on the grounds that—

- (1) the defendant is a biomaterials supplier; and
- (2)(A) the defendant should not, for the purposes of—
 - (i) section 205(b), be considered to be a manufacturer of the implant that is subject to such section; or
 - (ii) section 205(c), be considered to be a seller of the implant that allegedly caused harm to the claimant; or
- (B)(i) the claimant has failed to establish, pursuant to section 205(d), that the supplier furnished raw materials or component parts in violation of contractual requirements or specifications; or
- (ii) the claimant has failed to comply with the procedural requirements of subsection (b).

(b) **MANUFACTURER OF IMPLANT SHALL BE NAMED A PARTY.**—The claimant shall be required to name the manufacturer of the implant as a party to the action, unless—

- (1) the manufacturer is subject to service of process solely in a jurisdiction in which the biomaterials supplier is not domiciled or subject to a service of process; or
- (2) an action against the manufacturer is barred by applicable law.

(c) **PROCEEDING ON MOTION TO DISMISS.**—The following rules shall apply to any proceeding on a motion to dismiss filed under this section:

(1) **AFFIDAVITS RELATING TO LISTING AND DECLARATIONS.**—

(A) **IN GENERAL.**—The defendant in the action may submit an affidavit demonstrating that defendant has not included the implant on a list, if any, filed with the Secretary pursuant to section 510(j) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 360(j)).

(B) **RESPONSE TO MOTION TO DISMISS.**—In response to the motion to dismiss, the claimant may submit an affidavit demonstrating that—

(i) the Secretary has, with respect to the defendant and the implant that allegedly caused harm to the claimant, issued a declaration pursuant to section 205(b)(2)(B); or

(ii) the defendant who filed the motion to dismiss is a seller of the implant who is liable under section 205(c).

(2) **EFFECT OF MOTION TO DISMISS ON DISCOVERY.**—

(A) **IN GENERAL.**—If a defendant files a motion to dismiss under paragraph (1) or (2) of subsection (a), no discovery shall be permitted in connection to the action that is the subject of the motion, other than discovery necessary to determine a motion to dismiss for lack of jurisdiction, until such time as the court rules on the motion to dismiss in accordance with the affidavits submitted by the parties in accordance with this section.

(B) **DISCOVERY.**—If a defendant files a motion to dismiss under subsection (a)(2)(B)(i) on the grounds that the

biomaterials supplier did not furnish raw materials or component parts in violation of contractual requirements or specifications, the court may permit discovery, as ordered by the court. The discovery conducted pursuant to this subparagraph shall be limited to issues that are directly relevant to—

- (i) the pending motion to dismiss; or
- (ii) the jurisdiction of the court.

(3) AFFIDAVITS RELATING STATUS OF DEFENDANT.—

(A) IN GENERAL.—Except as provided in clauses (i) and (ii) of subparagraph (B), the court shall consider a defendant to be a biomaterials supplier who is not subject to an action for harm to a claimant caused by an implant, other than an action relating to liability for a violation of contractual requirements or specifications described in subsection (d).

(B) RESPONSES TO MOTION TO DISMISS.—The court shall grant a motion to dismiss any action that asserts liability of the defendant under subsection (b) or (c) of section 205 on the grounds that the defendant is not a manufacturer subject to such section 205(b) or seller subject to section 205(c), unless the claimant submits a valid affidavit that demonstrates that—

- (i) with respect to a motion to dismiss contending the defendant is not a manufacturer, the defendant meets the applicable requirements for liability as a manufacturer under section 205(b); or
- (ii) with respect to a motion to dismiss contending that the defendant is not a seller, the defendant meets the applicable requirements for liability as a seller under section 205(c).

(4) BASIS OF RULING ON MOTION TO DISMISS.—

(A) IN GENERAL.—The court shall rule on a motion to dismiss filed under subsection (a) solely on the basis of the pleadings of the parties made pursuant to this section and any affidavits submitted by the parties pursuant to this section.

(B) MOTION FOR SUMMARY JUDGMENT.—Notwithstanding any other provision of law, if the court determines that the pleadings and affidavits made by parties pursuant to this section raise genuine issues as concerning material facts with respect to a motion concerning contractual requirements and specifications, the court may deem the motion to dismiss to be a motion for summary judgment made pursuant to subsection (d).

(d) SUMMARY JUDGMENT.—

(1) IN GENERAL.—

(A) BASIS FOR ENTRY OF JUDGMENT.—A biomaterials supplier shall be entitled to entry of judgment without trial if the court finds there is no genuine issue as concerning any material fact for each applicable element set forth in paragraphs (1) and (2) of section 205(d).

(B) ISSUES OF MATERIAL FACT.—With respect to a finding made under subparagraph (A), the court shall consider a genuine issue of material fact to exist only if the evidence submitted by claimant would be sufficient to allow a reason-

able jury to reach a verdict for the claimant if the jury found the evidence to be credible.

(2) **DISCOVERY MADE PRIOR TO A RULING ON A MOTION FOR SUMMARY JUDGMENT.**—If, under applicable rules, the court permits discovery prior to a ruling on a motion for summary judgment made pursuant to this subsection, such discovery shall be limited solely to establishing whether a genuine issue of material fact exists as to the applicable elements set forth in paragraphs (1) and (2) of section 205(d).

(3) **DISCOVERY WITH RESPECT TO A BIOMATERIALS SUPPLIER.**—A biomaterials supplier shall be subject to discovery in connection with a motion seeking dismissal or summary judgment on the basis of the inapplicability of section 205(d) or the failure to establish the applicable elements of section 205(d) solely to the extent permitted by the applicable Federal or State rules for discovery against nonparties.

(e) **STAY PENDING PETITION FOR DECLARATION.**—If a claimant has filed a petition for a declaration pursuant to section 205(b)(3)(A) with respect to a defendant, and the Secretary has not issued a final decision on the petition, the court shall stay all proceedings with respect to that defendant until such time as the Secretary has issued a final decision on the petition.

(f) **MANUFACTURER CONDUCT OF PROCEEDING.**—The manufacturer of an implant that is the subject of an action covered under this title shall be permitted to file and conduct a proceeding on any motion for summary judgment or dismissal filed by a biomaterials supplier who is a defendant under this section if the manufacturer and any other defendant in such action enter into a valid and applicable contractual agreement under which the manufacturer agrees to bear the cost of such proceeding or to conduct such proceeding.

(g) **ATTORNEY FEES.**—The court shall require the claimant to compensate the biomaterials supplier (or a manufacturer appearing in lieu of a supplier pursuant to subsection (f)) for attorney fees and costs, if—

- (1) the claimant named or joined the biomaterials supplier; and
- (2) the court found the claim against the biomaterials supplier to be without merit and frivolous.

TITLE III—LIMITATIONS ON APPLICABILITY; EFFECTIVE DATE

SEC. 301. EFFECT OF COURT OF APPEALS DECISIONS.

A decision by a Federal circuit court of appeals interpreting a provision of this Act (except to the extent that the decision is overruled or otherwise modified by the Supreme Court) shall be considered a controlling precedent with respect to any subsequent decision made concerning the interpretation of such provision by any Federal or State court within the geographical boundaries of the area under the jurisdiction of the circuit court of appeals.

SEC. 302. FEDERAL CAUSE OF ACTION PRECLUDED.

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

SEC. 303. EFFECTIVE DATE.

This Act shall apply with respect to any action commenced on or after the date of the enactment of this Act without regard to whether the harm that is the subject of the action or the conduct that caused the harm occurred before such date of enactment.

BILL EMERSON,

Speaker of the House of Representatives pro tempore.

STROM THURMOND,

President of the Senate pro tempore.

[Endorsement on back of bill:]

I certify that this Act originated in the House of Representatives.

ROBIN H. CARLE, *Clerk.*

