CONSUMER PRODUCT SAFETY ACT

(Codified at 15 U.S.C. 2051–2084)

(Public Law 92-573; 86 Stat. 1207, Oct. 27, 1972)


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(References in brackets [ ] are to the United States Code and the Code of Federal Regulations)

(References in braces { } are editorial insertions)

[Public Law 92-573; Oct. 27, 1972, as amended]

SHORT TITLE; TABLE OF CONTENTS

SECTION 1. [15 U.S.C. 2051n] This Act may be cited as the “Consumer Product Safety Act”.

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FINDINGS AND PURPOSES

(a) The Congress finds that—

(1) an unacceptable number of consumer products which present unreasonable risks of injury are distributed in commerce;

(2) complexities of consumer products and the diverse nature and abilities of consumers using them frequently result in an inability of users to anticipate risks and to safeguard themselves adequately;

(3) the public should be protected against unreasonable risks of injury associated with consumer products;

(4) control by State and local governments of unreasonable risks of injury associated with consumer products is inadequate and may be burdensome to manufacturers;

(5) existing Federal authority to protect consumers from exposure to consumer products presenting unreasonable risks of injury is inadequate; and

(6) regulation of consumer products the distribution or use of which affects interstate or foreign commerce is necessary to carry out this Act.

(b) The purposes of this Act are—

(1) to protect the public against unreasonable risks of injury associated with consumer products;

(2) to assist consumers in evaluating the comparative safety of consumer products;

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(3) to develop uniform safety standards for consumer products and to minimize conflicting State and local regulations; and

(4) to promote research and investigation into the causes and prevention of product-related deaths, illnesses, and injuries.

DEFINITIONS

(a) For purposes of this Act:

(1) The term “consumer product” means any article, or component part thereof, produced or distributed (i) for sale to a consumer for use in or around a permanent or temporary household or residence, a school, in recreation, or otherwise, or (ii) for the personal use, consumption or enjoyment of a consumer in or around a permanent or temporary household or residence, a school, in recreation, or otherwise; but such term does not include—

(A) any article which is not customarily produced or distributed for sale to, or use or consumption by, or enjoyment of, a consumer,

(B) tobacco and tobacco products,

(C) motor vehicles or motor vehicle equipment (as defined by sections 102(3) and (4) of the National Traffic and Motor Vehicle Safety Act of 1966), [49 U.S.C. 30102(a)(6)-(7)]

(D) pesticides (as defined by the Federal Insecticide, Fungicide, and Rodenticide Act), [7 U.S.C. 136]

(E) any article which, if sold by the manufacturer, producer, or importer, would be subject to the tax imposed by section 4181 of the Internal Revenue Code of 1954 (determined without regard to any exemptions from such tax provided by section 4182 or 4221, or any other provision of such Code), or any component of any such article, firearms and ammunition

(F) aircraft, aircraft engines, propellers, or appliances (as defined in section 101 of the Federal Aviation Act of 1958), [49 U.S.C. 40102]

(G) boats which could be subjected to safety regulation under the Federal Boat Safety Act of 1971 (46 U.S.C. 1451 et seq.); [now codified as 46 U.S.C. 4301 et seq.] vessels, and appurtenances to vessels (other than such
boats), which could be subjected to safety regulation under title 52 of the Revised Statutes or other marine safety statutes administered by the department in which the Coast Guard is operating; and equipment (including associated equipment, as defined in section 3(8) of the Federal Boat Safety Act of 1971) [now codified as 46 U.S.C. 2101(1)] to the extent that a risk of injury associated with the use of such equipment on boats or vessels could be eliminated or reduced by action taken under any statute referred to in this subparagraph,

(H) drugs, devices, or cosmetics (as such terms are defined in sections 201 (g), (h), and (i) of the Federal Food, Drug, and Cosmetic Act), [21 U.S.C. 321(g), (h), (i)] or

(I) food. The term “food”, as used in this subparagraph means all “food”, as defined in section 201(f) of the Federal Food, Drug, and Cosmetic Act. [21 U.S.C. 321(f)] including poultry and poultry products (as defined in sections 4 (e) and (f) of the Poultry Products Inspection Act), [21 U.S.C. 453(e), (f)] meat, meat food products (as defined in section 1(j) of the Federal Meat Inspection Act), [21 U.S.C. 601(j)] and eggs and egg products (as defined in section 4 of the Egg Products Inspection Act). [21 U.S.C. 1033]

Such term includes any mechanical device which carries or conveys passengers along, around, or over a fixed or restricted route or course or within a defined area for the purpose of giving its passengers amusement, which is customarily controlled or directed by an individual who is employed for that purpose and who is not a consumer with respect to such device, and which is not permanently fixed to a site. Such term does not include such a device which is permanently fixed to a site. Except for the regulation under this Act or the Federal Hazardous Substances Act of fireworks devices or any substance intended for use as a component of any such device, the Commission shall have no authority under the functions transferred pursuant to section 30 of this Act to regulate any product or article described in subparagraph (E) of this paragraph or described, without regard to quantity, in section 845(a)(5) of title 18, United States Code. {Antique firearms supplies.} See sections 30(d) and 31 of
this Act, for other limitations on Commission’s authority to regulate certain consumer products.

(2) The term “consumer product safety rule” means a consumer products safety standard described in section 7(a), or a rule under this Act declaring a consumer product a banned hazardous product.

(3) The term “risk of injury” means a risk of death, personal injury, or serious or frequent illness.

(4) The term “manufacturer” means any person who manufactures or imports a consumer product.

(5) The term “distributor” means a person to whom a consumer product is delivered or sold for purposes of distribution in commerce, except that such term does not include a manufacturer or retailer of such product.

(6) The term “retailer” means a person to whom a consumer product is delivered or sold for purposes of sale or distribution by such person to a consumer.

(7)(A) The term “private labeler” means an owner of a brand or trademark on the label of a consumer product which bears a private label.

(B) A consumer product bears a private label if (i) the product (or its container) is labeled with the brand or trademark of a person other than a manufacturer of the product, (ii) the person with whose brand or trademark the product (or container) is labeled has authorized or caused the product to be so labeled, and (iii) the brand or trademark of a manufacturer of such product does not appear on such label.

(8) The term “manufactured” means to manufacture, produce, or assemble.


(10) The term “State” means a State, the District of Columbia, the Commonwealth of Puerto Rico, the Virgin Islands, Guam, Wake Island, Midway Island, Kingman Reef, Johnston Island, the Canal Zone, American Samoa, or the Trust Territory of the Pacific Islands.

(11) The terms “to distribute in commerce” and “distribution in commerce” means to sell in commerce, to introduce or deliver for introduction into commerce, or to hold for sale or distribution after introduction into commerce.

(12) The term “commerce” means trade, traffic, commerce, or transportation—

(A) between a place in a State and any place outside thereof, or
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(B) which affects trade, traffic, commerce, or transportation described in subparagraph (A).

(13) The terms “import” and “importation” include reimporting a consumer product manufactured or processed, in whole or in part, in the United States.

(14) The term “United States”, when used in the geographic sense, means all of the States (as defined in paragraph (10)).

(b) A common carrier, contract carrier, or freight forwarder shall not, for purposes of this Act, be deemed to be a manufacturer, distributor, or retailer of a consumer product solely by reason of receiving or transporting a consumer product in the ordinary course of its business as such a carrier or forwarder.

CONSUMER PRODUCT SAFETY COMMISSION


(a) An independent regulatory commission is hereby established, to be known as the Consumer Product Safety Commission, consisting of five Commissioners who shall be appointed by the President, by and with the advice and consent of the Senate. In making such appointments, the President shall consider individuals who, by reason of their background and expertise in areas related to consumer products and protection of the public from risks to safety, are qualified to serve as members of the Commission. The Chairman shall be appointed by the President, by and with the advice and consent of the Senate, from among the members of the Commission. An individual may be appointed as a member of the Commission and as Chairman at the same time. Any member of the Commission may be removed by the President for neglect of duty or malfeasance in office but for no other cause.

(b)(1) Except as provided in paragraph (2), (A) the Commissioners first appointed under this section shall be appointed for terms ending three, four, five, six, and seven years, respectively, after the date of the enactment of this Act, the term of each to be designated by the President at the time of nomination; and (B) each of their successors shall be appointed for a term of seven years from the date of the expiration of the term for which his predecessor was appointed.

(2) Any Commissioner appointed to fill a vacancy occurring prior to the expiration of the term for which his predecessor was appointed shall be appointed only for the remainder of such term. A Commissioner may continue to serve after the expiration of his term until his successor has taken office,

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except that he may not so continue to serve more than one year after the date on which his term would otherwise expire under this subsection.

(c) Not more than three of the Commissioners shall be affiliated with the same political party. No individual (1) in the employ of, or holding any official relation to, any person engaged in selling or manufacturing consumer products, or (2) owning stock or bonds of substantial value in a person so engaged, or (3) who is in any other manner pecuniarily interested in such a person, or in a substantial supplier of such a person, shall hold the office of Commissioner. A Commissioner may not engage in any other business, vocation, or employment.

(d) No vacancy in the Commission shall impair the right of the remaining Commissioners to exercise all the powers of the Commission, but three members of the Commission shall constitute a quorum for the transaction of business, except that if there are only three members serving on the Commission because of vacancies in the Commission, two members of the Commission shall constitute a quorum for the transaction of business, and if there are only two members serving on the Commission because of vacancies in the Commission, two members shall constitute a quorum for the six month period beginning on the date of the vacancy which caused the number of Commission members to decline to two. The Commission shall have an official seal of which judicial notice shall be taken. The Commission shall annually elect a Vice Chairman to act in the absence or disability of the Chairman or in case of a vacancy in the office of the Chairman.

(e) The Commission shall maintain a principal office and such field offices as it deems necessary and may meet and exercise any of its powers at any other place.

(f)(1) The Chairman of the Commission shall be the principal executive officer of the Commission, and he shall exercise all of the executive and administrative functions of the Commission, including functions of the Commission with respect to (A) the appointment and supervision of personnel employed under the Commission (other than personnel employed regularly and full time in the immediate offices of commissioners other than the Chairman), (B) the distribution of business among personnel appointed and supervised by the Chairman and among administrative units of the Commission, and (C) the use and expenditure of funds.

(2) In carrying out any of his functions under the provisions of this subsection the Chairman shall be governed by general

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policies of the Commission and by such regulatory decisions, findings, and determinations as the Commission may by law be authorized to make.

(3) Requests or estimates for regular, supplemental, or deficiency appropriations on behalf of the Commission may not be submitted by the Chairman without the prior approval of the Commission.

(g)(1)(A) The Chairman, subject to the approval of the Commission, shall appoint as officers of the Commission an Executive Director, a General Counsel, an Associate Executive Director for Engineering Sciences, an Associate Executive Director for Epidemiology, an Associate Executive Director for Compliance and Administrative Litigation, an Associate Executive Director for Health Sciences, an Associate Executive Director for Economic Analysis, an Associate Executive Director for Administration, an Associate Executive Director for Field Operations, a Director for Office of Program, Management, and Budget, and a Director for Office of Information and Public Affairs. Any other individual appointed to a position designated as an Associate Executive Director shall be appointed by the Chairman, subject to the approval of the Commission. The Chairman may only appoint an attorney to the position of Associate Executive Director of Compliance and Administrative Litigation except the position of acting Associate Executive Director of Compliance and Administrative Litigation.

(B)(i) No individual may be appointed to such a position on an acting basis for a period longer than 90 days unless such appointment is approved by the Commission.

(ii) The Chairman, with the approval of the Commission, may remove any individual serving in a position appointed under subparagraph (A).

(C) Subparagraph (A) shall not be construed to prohibit appropriate reorganizations or changes in classification.

(2) The Chairman, subject to subsection (f)(2), may employ such other officers and employees (including attorneys) as are necessary in the execution of the Commission’s functions.

(3) In addition to the number of positions authorized by section 5108(a) of title 5, United States Code, the Chairman, subject to the approval of the Commission, and subject to the standards and procedures prescribed by chapter 51 of title 5, United States Code, may place a total of twelve positions in grades GS-16, GS-17, and GS-18.

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an agency (as defined in section 5102(a)(1) of such title 5) under any such provision to place one or more positions in GS-16, 17, or 18 of the General Schedule, is hereby terminated.”}

(4) The appointment of any officer (other than a Commissioner) or employee of the Commission shall not be subject, directly or indirectly, to review or approval by any officer or entity within the Executive Office of the President.

(h) (i) Subsections (a) and (h) of section 2680 of title 28, United States Code, do not prohibit the bringing of a civil action on a claim against the United States which—

(1) is based upon—

(A) misrepresentation or deceit on the part of the Commission or any employee thereof, or

(B) any exercise or performance, or failure to exercise or perform, a discretionary function on the part of the Commission or any employee thereof which exercise, performance, or failure was grossly negligent; and

(2) is not made with respect to any agency action (as defined in section 551(13) of title 5, United States Code).

In the case of a civil action on a claim based upon the exercise or performance of, or failure to exercise or perform, a discretionary function, no judgment may be entered against the United States unless the court in which such action was brought determines (based upon consideration of all the relevant circumstances, including the statutory responsibility of the Commission and the public interest in encouraging rather than inhibiting the exercise of discretion) that such exercise, performance, or failure to exercise or perform was unreasonable.

(j) At least 30 days before the beginning of each fiscal year, the Commission shall establish an agenda for Commission action under the Acts under its jurisdiction and, to the extent feasible, shall establish priorities for such actions. Before establishing such agenda and priorities, the Commission shall conduct a public hearing on the agenda and priorities and shall provide reasonable opportunity for the submission of comments. {This subsection effective FY 92.}

PRODUCT SAFETY INFORMATION AND RESEARCH


(a) The Commission shall—

(1) maintain an Injury Information Clearinghouse to collect, investigate, analyze, and disseminate injury

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data, and information, relating to the causes and prevention of death, injury, and illness associated with consumer products;

(2) conduct such continuing studies and investigations of deaths, injuries, diseases, other health impairments, and economic losses resulting from accidents involving consumer products as it deems necessary;

(3) following publication of an advance notice of proposed rulemaking or a notice of proposed rulemaking for a product safety rule under any rulemaking authority administered by the Commission, assist public and private organizations or groups of manufacturers, administratively and technically, in the development of safety standards addressing the risk of injury identified in such notice; and

(4) to the extent practicable and appropriate (taking into account the resources and priorities of the Commission), assist public and private organizations or groups of manufacturers, administratively and technically, in the development of product safety standards and test methods.

(b) The Commission may—

(1) conduct research, studies, and investigations on the safety of consumer products and on improving the safety of such products;

(2) test consumer products and develop product safety test methods and testing devices; and

(3) offer training in product safety investigation and test methods.

(c) In carrying out its functions under this section, the Commission may make grants or enter into contracts for the conduct of such functions with any person (including a governmental entity).

(d) Whenever the Federal contribution for any information, research, or development activity authorized by this Act is more than minimal, the Commission shall include in any contract, grant, or other arrangement for such activity, provisions effective to insure that the rights to all information, uses, processes, patents, and other developments resulting from that activity will be made available to the public without charge on a nonexclusive basis. Nothing in this subsection shall be construed to deprive any person of any right which he may have had, prior to entering into any arrangement referred to in this subsection, to any patent, patent application, or invention.

\*Note: 35 U.S.C. 200-211 and 37 CFR Part 401 specifically supersede section 5(d) of the Consumer Product Safety Act with respect to small

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business firms and nonprofit organizations which retain, in most cases, exclusive commercial rights to inventions made with Commission support.

PUBLIC DISCLOSURE OF INFORMATION


(a)(1) Nothing contained in this Act shall be construed to require the release of any information described by subsection (b) of section 552 of title 5, United States Code, or which is otherwise protected by law from disclosure to the public.

(2) All information reported to or otherwise obtained by the Commission or its representative under this Act which information contains or relates to a trade secret or other matter referred to in section 1905 of title 18, United States Code, or subject to section 552(b)(4) of title 5, United States Code, shall be considered confidential and shall not be disclosed.

(3) The Commission shall, prior to the disclosure of any information which will permit the public to ascertain readily the identity of a manufacturer or private labeler of a consumer product, offer such manufacturer or private labeler an opportunity to mark such information as confidential and therefore barred from disclosure under paragraph (2).

(4) All information that a manufacturer or private labeler has marked to be confidential and barred from disclosure under paragraph (2), either at the time of submission or pursuant to paragraph (3), shall not be disclosed, except in accordance with the procedures established in paragraphs (5) and (6).

(5) If the Commission determines that a document marked as confidential by a manufacturer or private labeler to be barred from disclosure under paragraph (2) may be disclosed because it is not confidential information as provided in paragraph (2), the Commission shall notify such person in writing that the Commission intends to disclose such document at a date not less than 10 days after the date of receipt of notification.

(6) Any person receiving such notification may, if he believes such disclosure is barred by paragraph (2), before the date set for release of the document, bring an action in the district court of the United States in the district in which the complainant resides, or has his principal place of business, or in which the documents are located, or in the United States District Court for the District of Columbia to restrain disclosure of the document. Any person receiving such notification may file with the appropriate district court or court of appeals of the United States, as appropriate, an application for a stay of disclosure. The documents shall not be disclosed until the court has ruled on the application for a stay.
(7) Nothing in this Act shall authorize the withholding of information by the Commission or any officer or employee under its control from the duly authorized committees or subcommittees of the Congress, and the provisions of paragraphs (2) through (6) shall not apply to such disclosures, except that the Commission shall immediately notify the manufacturer or private labeler of any such request for information designated as confidential by the manufacturer or private labeler.

(8) The provisions of paragraphs (2) through (6) shall not prohibit the disclosure of information to other officers, employees, or representatives of the Commission (including contractors) concerned with carrying out this Act or when relevant in any administrative proceeding under this Act or in judicial proceedings to which the Commission is a party. Any disclosure of relevant information—

(A) in Commission administrative proceedings or in judicial proceedings to which the Commission is a party, or

(B) to representatives of the Commission (including contractors),

shall be governed by the rules of the Commission (including in camera review rules for confidential material) for such proceedings or for disclosures to such representatives or by court rules or orders, except that the rules of the Commission shall not be amended in a manner inconsistent with the purpose of this section.

(b)(1) Except as provided by paragraph (4) of this subsection, not less than 30 days prior to its public disclosure of any information obtained under this Act, or to be disclosed to the public in connection therewith (unless the Commission finds that the public health and safety requires a lesser period of notice and publishes such a finding in the Federal Register), the Commission shall, to the extent practicable, notify and provide a summary of the information to, each manufacturer or private labeler of any consumer product to which such information pertains, if the manner in which such consumer product is to be designated or described in such information will permit the public to ascertain readily the identity of such manufacturer or private labeler, and shall provide such manufacturer or private labeler with a reasonable opportunity to submit comments to the Commission in regard to such information. The Commission shall take reasonable steps to assure, prior to its public disclosure thereof, that information from which the identity of such manufacturer or private labeler may be readily ascertained is accurate, and that such disclosure is fair in the circumstances and reasonably related to

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effectuating the purposes of this Act. In disclosing any information under this subsection, the Commission may, and upon the request of the manufacturer or private labeler shall, include with the disclosure any comments or other information or a summary thereof submitted by such manufacturer or private labeler to the extent permitted by and subject to the requirements of this section.

(2) If the Commission determines that a document claimed to be inaccurate by a manufacturer or private labeler under paragraph (1) should be disclosed because the Commission believes it has complied with paragraph (1), the Commission shall notify the manufacturer or private labeler that the Commission intends to disclose such document at a date not less than 10 days after the date of the receipt of notification. The Commission may provide a lesser period of notice of intent to disclose if the Commission finds that the public health and safety requires a lesser period of notice and publishes such finding in the Federal Register.

(3) Prior to the date set for release of the document, the manufacturer or private labeler receiving the notice described in paragraph (2) may bring an action in the district court of the United States in the district in which the complainant resides, or has his principal place of business, or in which the documents are located or in the United States District Court for the District of Columbia to enjoin disclosure of the document. The district court may enjoin such disclosure if the Commission has failed to take the reasonable steps prescribed in paragraph (1).

(4) Paragraphs (1) through (3) of this subsection shall not apply to the public disclosure of (A) information about any consumer product with respect to which product the Commission has filed an action under section 12 (relating to imminently hazardous products), or which the Commission has reasonable cause to believe is in violation of section 19 (relating to prohibited acts); or (B) information in the course of or concerning a rulemaking proceeding (which shall commence upon the publication of an advance notice of proposed rulemaking or a notice of proposed rulemaking), an adjudicatory proceeding (which shall commence upon the issuance of a complaint) or other administrative or judicial proceeding under this Act.

(5) In addition to the requirements of paragraph (1), the Commission shall not disclose to the public information submitted pursuant to section 15(b) respecting a consumer product unless—

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(A) the Commission has issued a complaint under section 15 (c) or (d) alleging that such product presents a substantial product hazard;

(B) in lieu of proceeding against such product under section 15 (c) or (d), the Commission has accepted in writing a remedial settlement agreement dealing with such product; or

(C) the person who submitted the information under section 15(b) agrees to its public disclosure.

The provisions of this paragraph shall not apply to the public disclosure of information with respect to a consumer product which is the subject of an action brought under section 12, or which the Commission has reasonable cause to believe is in violation of section 19(a), or information in the course of or concerning a judicial proceeding.

(6) Where the Commission initiates the public disclosure of information that reflects on the safety of a consumer product or class of consumer products, whether or not such information would enable the public to ascertain readily the identity of a manufacturer or private labeler, the Commission shall establish procedures designed to ensure that such information is accurate and not misleading.

(7) If the Commission finds that, in the administration of this Act, it has made public disclosure of inaccurate or misleading information which reflects adversely upon the safety of any consumer product or class of consumer products, or the practices of any manufacturer, private labeler, distributor, or retailer of consumer products, it shall, in a manner equivalent to that in which such disclosure was made, take reasonable steps to publish a retraction of such inaccurate or misleading information.

(8) If, after the commencement of a rulemaking or the initiation of an adjudicatory proceeding, the Commission decides to terminate the proceeding before taking final action, the Commission shall, in a manner equivalent to that in which such commencement or initiation was publicized, take reasonable steps to make known the decision to terminate.

(c) The Commission shall communicate to each manufacturer of a consumer product, insofar as may be practicable, information as to any significant risk of injury associated with such product.

(d)(1) For purposes of this section, the term “Act” means the Consumer Product Safety Act, the Flammable Fabrics Act, the Poison Prevention Packaging Act, and the Federal Hazardous Substances Act.
(2) The provisions of this section shall apply whenever information is to be disclosed by the Commission, any member of the Commission, or any employee, agent, or representative of the Commission in an official capacity.

(e)(1) Notwithstanding the provisions of section 552 of title 5, United States Code, subsection (a)(7) of this section, or of any other law, except as provided in paragraphs (2), (3), and (4), no member of the Commission, no officer or employee of the Commission, and no officer or employee of the Department of Justice may—

(A) publicly disclose information furnished under subsection (c)(1) or (c)(2)(A) of section 37;

(B) use such information for any purpose other than to carry out the Commission’s responsibilities; or

(C) permit anyone (other than the members, officers, and employees of the Commission or officers or employees of the Department of Justice who require such information for an action filed on behalf of the Commission) to examine such information.

(2) Any report furnished under subsection (c)(1) or (c)(2)(A) of section 37 shall be immune from legal process and shall not be subject to subpoena or other discovery in any civil action in a State or Federal court or in any administrative proceeding, except in an action against such manufacturer under section 20, 21, or 22 for failure to furnish information required by section 37.

(3) The Commission may, upon written request, furnish to any manufacturer or to the authorized agent of such manufacturer authenticated copies of reports furnished by or on behalf of such manufacturer in accordance with section 37, upon payment of the actual or estimated cost of searching the records and furnishing such copies.

(4) Upon written request of the Chairman or Ranking Minority Member of the Committee on Commerce, Science, and Transportation of the Senate or the Committee on Energy and Commerce of the House of Representatives or any subcommittee of such committee, the Commission shall provide to the Chairman or Ranking Minority Member any information furnished to the Commission under section 37 for purposes that are related to the jurisdiction of such committee or subcommittee.

(5) Any officer or employee of the Commission or other officer or employee of the Federal Government who receives information provided under section 37, who willfully violates the requirements of this subsection shall be subject to dismissal or

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other action consistent with procedures and requirements established by the Office of Personnel Management.

CONSUMER PRODUCT SAFETY STANDARDS


(a) The Commission may promulgate consumer product safety standards in accordance with the provisions of section 9. A consumer product safety standard shall consist of one or more of any of the following types of requirements:

(1) Requirements expressed in terms of performance requirements.

(2) Requirements that a consumer product be marked with or accompanied by clear and adequate warnings or instructions, or requirements respecting the form of warnings or instructions.

Any requirement of such a standard shall be reasonably necessary to prevent or reduce an unreasonable risk of injury associated with such product.

(b)(1) The Commission shall rely upon voluntary consumer product safety standards rather than promulgate a consumer product safety standard prescribing requirements described in subsection (a) whenever compliance with such voluntary standards would eliminate or adequately reduce the risk of injury addressed and it is likely that there will be substantial compliance with such voluntary standards.

(2) The Commission shall devise procedures to monitor compliance with any voluntary standards—.

(A) upon which the Commission has relied under paragraph (1);

(B) which were developed with the participation of the Commission; or

(C) whose development the Commission has monitored.

(c) If any person participates with the Commission in the development of a consumer product safety standard, the Commission may agree to contribute to the person’s cost with respect to such participation, in any case in which the Commission determines that such contribution is likely to result in a more satisfactory standard than would be developed without such contribution, and that the person is financially responsible. Regulations of the Commission [16 CFR Part 1105] shall set forth the items of cost in which it may participate, and shall exclude any contribution to the acquisition of land or buildings. Payments under agreements entered into under this subsection may be made without regard to section 3648 of the

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BANNED HAZARDOUS PRODUCTS

Whenever the Commission finds that—

(1) a consumer product is being, or will be, distributed in commerce and such consumer product presents an unreasonable risk of injury; and

(2) no feasible consumer product safety standard under this Act would adequately protect the public from the unreasonable risk of injury associated with such product,

the Commission may, in accordance with section 9, promulgate a rule declaring such product a banned hazardous product.

BANNING OF BUTYL NITRITE

[Sec. 2404 of Pub. L. 100-690; 15 U.S.C. 2057a]
\{Not technically part of the Consumer Product Safety Act\}

(a) IN GENERAL.—Except as provided in subsection (b), butyl nitrite shall be considered a banned hazardous product under section 8 of the Consumer Product Safety Act (15 U.S.C. 2057).

(b) LAWFUL PURPOSES.—For the purposes of section 8 of the Consumer Product Safety Act, it shall not be unlawful for any person to manufacture for sale, offer for sale, distribute in commerce, or import into the United States butyl nitrite for any commercial purpose or any other purpose approved under the Federal Food, Drug, and Cosmetic Act.

(c) DEFINITIONS.—For purposes of this section:

(1) The term “butyl nitrite” includes n-butyl nitrite, isobutyl nitrite, secondary butyl nitrite, tertiary butyl nitrite, and mixtures containing these chemicals.

(2) The term “commercial purpose” means any commercial purpose other than for the production of consumer products containing butyl nitrite that may be used for inhaling or otherwise introducing butyl nitrite into the human body for euphoric or physical effects.

(d) EFFECTIVE DATE.—This section shall take effect 90 days after the date of the enactment of this subtitle. {enacted November 18, 1988}
BANNING OF ISOPROPAL NITRITE AND OTHER NITRITES

[Title XXIII, Sec. 3202 of Pub. L. 101-647; 15 U.S.C. 2057b]

Not technically part of the Consumer Product Safety Act

(a) IN GENERAL.—Except as provided in subsection (b), volatile alkyl nitrite shall be considered a banned hazardous product under section 8 of the Consumer Product Safety Act (15 U.S.C. 2057).

(b) LAWFUL PURPOSES.—For the purposes of section 8 of the Consumer Product Safety Act, it shall not be unlawful for any person to manufacture for sale, offer for sale, distribute in commerce, or import into the United States volatile alkyl nitrites for any commercial purpose or any other purpose approved under the Federal Food, Drug, and Cosmetic Act.

(c) DEFINITIONS.—For purposes of this section, the term “commercial purpose” means any commercial purpose other than for the production of consumer products containing volatile alkyl nitrites that may be used for inhaling or otherwise introducing volatile alkyl nitrites into the human body for euphoric or physical effects.

(d) EFFECTIVE DATE.—This section shall take effect 90 days after the date of the enactment of this Act. [enacted November 29, 1990]

PROCEDURE FOR CONSUMER PRODUCT SAFETY RULES


(a) A proceeding for the development of a consumer product safety rule shall be commenced by the publication in the Federal Register of an advance notice of proposed rulemaking which shall—

1) identify the product and the nature of the risk of injury associated with the product;

2) include a summary of each of the regulatory alternatives under consideration by the Commission (including voluntary consumer product safety standards);

3) include information with respect to any existing standard known to the Commission which may be relevant to the proceedings, together with a summary of the reasons why the Commission believes preliminarily that such standard does not eliminate or adequately reduce the risk of injury identified in paragraph (1);

4) invite interested persons to submit to the Commission, within such period as the Commission shall specify in the notice (which period shall not be
CONSUMER PRODUCT SAFETY ACT

less than 30 days or more than 60 days after the date of publication of the notice), comments with respect to the risk of injury identified by the Commission, the regulatory alternatives being considered, and other possible alternatives for addressing the risk;

(5) invite any person (other than the Commission) to submit to the Commission, within such period as the Commission shall specify in the notice (which period shall not be less than 30 days after the date of publication of the notice), an existing standard or a portion of a standard as a proposed consumer product safety standard; and

(6) invite any person (other than the Commission) to submit to the Commission, within such period as the Commission shall specify in the notice (which period shall not be less than 30 days after the date of publication of the notice), a statement of intention to modify or develop a voluntary consumer product safety standard to address the risk of injury identified in paragraph (1) together with a description of a plan to modify or develop the standard.

The Commission shall transmit such notice within 10 calendar days to the Committee on Commerce, Science, and Transportation of the Senate and the Committee on Energy and Commerce of the House of Representatives. {This reporting requirement ceased to be effective on December 21, 1999 per Pub. L. 104-66, § 3003.}

(b)(1) If the Commission determines that any standard submitted to it in response to an invitation in a notice published under subsection (a)(5) if promulgated (in whole, in part, or in combination with any other standard submitted to the Commission or any part of such a standard) as a consumer product safety standard, would eliminate or adequately reduce the risk of the injury identified in the notice under subsection (a)(1), the Commission may publish such standard, in whole, in part, or in such combination and with nonmaterial modifications, as proposed consumer product safety rule.

(2) If the Commission determines that—

(A) compliance with any standard submitted to it in response to an invitation in a notice published under subsection (a)(6) is likely to result in the elimination or adequate reduction of the risk of injury identified in the notice, and

(B) it is likely that there will be substantial compliance with such standard,

the Commission shall terminate any proceeding to promulgate a consumer product safety rule respecting such risk of injury and

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shall publish in the Federal Register a notice which includes the
determination of the Commission and which notifies the public
that the Commission will rely on the voluntary standard to
eliminate or reduce the risk of injury, except that the
Commission shall terminate any such proceeding and rely on a
voluntary standard only if such voluntary standard is in
existence. For purposes of this section, a voluntary standard
shall be considered to be in existence when it is finally approved
by the organization or other person which developed such
standard, irrespective of the effective date of the standard.
Before relying upon any voluntary consumer product safety
standard, the Commission shall afford interested persons
(including manufacturers, consumers, and consumer
organizations) a reasonable opportunity to submit written
comments regarding such standard. The Commission shall
consider such comments in making any determination
regarding reliance on the involved voluntary standard under
this subsection.

(c) No consumer product safety rule may be proposed by the
Commission unless, not less than 60 days after publication of
the notice required in subsection (a), the Commission publishes
in the Federal Register the text of the proposed rule, including
any alternatives, which the Commission proposes to
promulgate, together with a preliminary regulatory analysis
containing—

(1) a preliminary description of the potential
benefits and potential costs of the proposed rule,
including any benefits or costs that cannot be
quantified in monetary terms, and an identification of
those likely to receive the benefits and bear the costs;
(2) a discussion of the reasons any standard or
portion of a standard submitted to the Commission
under subsection (a)(5) was not published by the
Commission as the proposed rule or part of the
proposed rule;
(3) a discussion of the reasons for the Commission’s
preliminary determination that efforts proposed under
subsection (a)(6) and assisted by the Commission as
required by section 5(a)(3) would not, within a
reasonable period of time, be likely to result in the
development of a voluntary consumer product safety
standard that would eliminate or adequately reduce the
risk of injury addressed by the proposed rule; and
(4) a description of any reasonable alternatives to
the proposed rule, together with a summary description
of their potential costs and benefits, and a brief
explanation of why such alternatives should not be
published as a proposed rule.
The Commission shall transmit such notice within 10 calendar days to the Committee on Commerce, Science, and Transportation of the Senate and the Committee on Energy and Commerce {now Committee on Commerce} of the House of Representatives. {This reporting requirement ceased to be effective on December 21, 1999 per Pub. L. 104-66, § 3003.} Any proposed consumer product safety rule shall be issued within twelve months after the date of publication of an advance notice of proposed rulemaking under subsection (a) relating to the product involved, unless the Commission determines that such proposed rule is not reasonably necessary to eliminate or reduce the risk of injury associated with the product or is not in the public interest. The Commission may extend the twelve-month period for good cause. If the Commission extends such period, it shall immediately transmit notice of such extension to the Committee on Commerce, Science, and Transportation of the Senate and the Committee on Energy and Commerce {now Committee on Commerce} of the House of Representatives. Such notice shall include an explanation of the reasons for such extension, together with an estimate of the date by which the Commission anticipates such rulemaking will be completed. The Commission shall publish notice of such extension and the information submitted to the Congress in the Federal Register.

(d)(1) Within 60 days after the publication under subsection (c) of a proposed consumer product safety rule respecting a risk of injury associated with a consumer product, the Commission shall—

(A) promulgate a consumer product safety rule respecting the risk of injury associated with such product, if it makes the findings required under subsection (f), or

(B) withdraw the applicable notice of proposed rulemaking if it determines that such rule is not (1) reasonably necessary to eliminate or reduce an unreasonable risk of injury associated with the product, or (ii) in the public interest;

except that the Commission may extend such 60-day period for good cause shown (if it publishes its reasons therefor in the Federal Register).

(2) Consumer product safety rules shall be promulgated in accordance with section 553 of title 5, United States Code, except that the Commission shall give interested persons an opportunity for the oral presentation of data, views, or arguments, in addition to an opportunity to make written submissions. A transcript shall be kept of any oral presentation.

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(e) A consumer product safety rule shall express in the rule itself the risk of injury which the standard is designed to eliminate or reduce. In promulgating such a rule the Commission shall consider relevant available product data including the results of research, development, testing, and investigation activities conducted generally and pursuant to this Act. In the promulgation of such a rule the Commission shall also consider and take into account the special needs of elderly and handicapped persons to determine the extent to which such persons may be adversely affected by such rule.

(f)(1) Prior to promulgating a consumer product safety rule, the Commission shall consider, and shall make appropriate findings for inclusion in such rule with respect to—

(A) the degree and nature of the risk of injury the rule is designed to eliminate or reduce;
(B) the approximate number of consumer products, or types or classes thereof, subject to such rule;
(C) the need of the public for the consumer products subject to such rule, and the probable effect of such rule upon the utility, cost, or availability of such products to meet such need; and
(D) any means of achieving the objective of the order while minimizing adverse effects on competition or disruption or dislocation of manufacturing and other commercial practices consistent with the public health and safety.

(2) The Commission shall not promulgate a consumer product safety rule unless it has prepared, on the basis of the findings of the Commission under paragraph (1) and on other information before the Commission, a final regulatory analysis of the rule containing the following information:

(A) A description of the potential benefits and potential costs of the rule, including costs and benefits that cannot be quantified in monetary terms, and the identification of those likely to receive the benefits and bear the costs.
(B) A description of any alternatives to the final rule which were considered by the Commission, together with a summary description of their potential benefits and costs and a brief explanation of the reasons why these alternatives were not chosen.
(C) A summary of any significant issues raised by the comments submitted during the public comment period in response to the preliminary regulatory analysis, and a summary of the assessment by the Commission of such issues.
The Commission shall publish its final regulatory analysis with the rule.

(3) The Commission shall not promulgate a consumer product safety rule unless it finds (and includes such finding in the rule)—

(A) that the rule (including its effective date) is reasonably necessary to eliminate or reduce an unreasonable risk of injury associated with such product;

(B) that the promulgation of the rule is in the public interest;

(C) in the case of a rule declaring the product a banned hazardous product, that no feasible consumer product safety standard under this Act would adequately protect the public from the unreasonable risk of injury associated with such product;

(D) in the case of a rule which relates to a risk of injury with respect to which persons who would be subject to such rule have adopted and implemented a voluntary consumer product safety standard, that—

(i) compliance with such voluntary consumer product safety standard is not likely to result in the elimination or adequate reduction of such risk of injury; or

(ii) it is unlikely that there will be substantial compliance with such voluntary consumer product safety standard;

(E) that the benefits expected from the rule bear a reasonable relationship to its costs; and

(F) that the rule imposes the least burdensome requirement which prevents or adequately reduces the risk of injury for which the rule is being promulgated.

(4)(A) Any preliminary or final regulatory analysis prepared under subsection (c) or (f)(2) shall not be subject to independent judicial review, except that when an action for judicial review of a rule is instituted, the contents of any such regulatory analysis shall constitute part of the whole rulemaking record of agency action in connection with such review.

(B) The provisions of subparagraph (A) shall not be construed to alter the substantive or procedural standards otherwise applicable to judicial review of any action by the Commission.

(g)(1) Each consumer product safety rule shall specify the date such rule is to take effect not exceeding 180 days from the date promulgated, unless the Commission finds, for good cause shown, that a later effective date is in the public interest and publishes its reasons for such finding. The effective date of a
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consumer product safety standard under this Act shall be set at a date at least 30 days after the date of promulgation unless the Commission for good cause shown determines that an earlier effective date is in the public interest. In no case may the effective date be set at a date which is earlier than the date of promulgation. A consumer product safety standard shall be applicable only to consumer products manufactured after the effective date.

(2) The Commission may by rule prohibit a manufacturer of a consumer product from stockpiling any product to which a consumer product safety rule applies, so as to prevent such manufacturer from circumventing the purpose of such consumer product safety rule. For purposes of this paragraph, the term “stockpiling” means manufacturing or importing a product between the date of promulgation of such consumer product safety rule and its effective date at a rate which is significantly greater (as determined under the rule under this paragraph) than the rate at which such product was produced or imported during a base period (prescribed in the rule under this paragraph) ending before the date of promulgation of the consumer product safety rule.

(h) The Commission may by rule amend or revoke any consumer product safety rule. Such amendment or revocation shall specify the date on which it is to take effect which shall not exceed 180 days from the date the amendment or revocation is published unless the Commission finds for good cause shown that a later effective date is in the public interest and publishes its reasons for such finding. Where an amendment involves a material change in a consumer product safety rule, sections 7 and 8, and subsections (a) through (g) of this section shall apply. In order to revoke a consumer product safety rule, the Commission shall publish a proposal to revoke such rule in the Federal Register, and allow oral and written presentations in accordance with subsection (d)(2) of this section. It may revoke such rule only if it determines that the rule is not reasonably necessary to eliminate or reduce an unreasonable risk of injury associated with the product. Section 11 shall apply to any amendment of a consumer product safety rule which involves a material change and to any revocation of a consumer product safety rule, in the same manner and to the same extent as such section applies to the Commission’s action in promulgating such a rule.

(i) The Commission shall grant, in whole or in part, or deny any petition under section 553(e) of Title 5, United State Code, requesting the Commission to initiate a rulemaking, within a reasonable time after the date on which such petition is filed. The Commission shall state the reasons for granting or denying

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such petition. The Commission may not deny any such petition on the basis of a voluntary standard unless the voluntary standard is in existence at the time of the denial of the petition, the Commission has determined that the voluntary standard is likely to result in the elimination or adequate reduction of the risk of injury identified in the petition, and it is likely that there will be substantial compliance with the standard.

LAWN MOWER STANDARD AMENDMENT

\{Not technically part of the Consumer Product Safety Act\}

(a) Not later than 90 days after the date of this Act, the Consumer Product Safety Commission shall amend its consumer product safety standard for walk-behind power lawn mowers to provide that a manually started rotary type lawn mower which has a blade control system which meets the requirements of the standard relating to blade controls (16 CFR 1205.5) except that the system stops the engine and requires a manual restart of the engine shall be considered in compliance with such requirements if the engine starting controls for the lawn mower are located within twenty-four inches from the top of the mower’s handles or the mower has a protective foot shield which extends three hundred and sixty degrees around the mower housing. The Consumer Product Safety Act shall not apply with respect to the promulgation of the amendment prescribed by this subsection.

(b) The Commission shall conduct a study of the effect on consumers of the amendment prescribed by subsection (a) and shall report the results of such study two years after the date the standard, as amended in accordance with subsection (a), takes effect. The Commission may not amend the amendment prescribed by subsection (a) before the report is filed under this subsection.

LAWN DARTS

[Public Law 100-613, 102 Stat. 3183, November 5, 1989]
\{Not technically part of the Consumer Product Safety Act\}

An Act

To provide that the Consumer Product Safety Commission amend its regulations regarding lawn darts.

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

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provision of law, not later that 60 days after the date of enactment of this Act. \footnote{enacted November 5, 1988} the Consumer Product Safety Commission shall amend its regulations to revoke exemption regarding lawn darts and other similar sharp-pointed toys contained in section 1500.86(a)(3) of title 16, Code of Federal Regulations, unless the Commission finds that such products do not have the potential for causing puncture wound injury.

AUTOMATIC GARAGE DOOR OPENERS


(a) Consumer Product Safety Rule.—The provision of subsection (b) shall be considered to be a consumer product safety rule issued by the Consumer Product Safety Commission under section 9 of the Consumer Product Safety Act.

(b) Requirements.—


2. Effective on and after January 1, 1993, all residential automatic garage door openers manufactured on and after such date for sale in the United States shall conform to any additional entrapment protection requirements of the American National Standards Institute Underwriters Laboratories, Inc. Standards for Safety—UL 325, third edition, which were issued after the date of the enactment of this Act to become effective on or before January 1, 1993.

B If, by June 1, 1992, the Underwriters Laboratories, Inc. has not issued a revision to the May 4, 1988, Standards for Safety—UL 325, third edition, to require an entrapment protection feature or device in addition to that required by the May 4, 1988, Standard, the Consumer Product Safety Commission shall begin a rulemaking proceeding, to be completed no later than October 31, 1992, to require an additional such feature or device on all automatic residential garage door openers manufactured on or after January 1, 1993, for sale in the United States. If such a revision is issued by the Underwriters Laboratories, Inc. after the rulemaking has commenced, the rulemaking shall be
terminated and the revision shall be incorporated in the consumer product safety rule under subsection (a) unless the Commission has determined under subsection (c) that such revision does not carry out the purposes of subsection (b).

(c) Revision of Rule.—If, after June 1, 1992, or the date of revision described in subsection (b)(2)(B) if later, the Underwriters Laboratories, Inc proposes to further revise the entrapment protection requirements of the American National Standards Institute Underwriters Laboratories, Inc. Standards for Safety—UL 325, third edition, the Laboratories shall notify the Consumer Product Safety Commission of the proposed revision and the proposed revision shall be incorporated in the consumer product safety rule under subsection (a) unless, within 30 days of such notice, the Commission notifies the Laboratories that the Commission has determined that such revision does not carry out the purposes of subsection (b).

(d) Labeling.—On and after January 1, 1991, a manufacturer selling or offering for sale in the United States an automatic residential garage door opener manufactured on or after January 1, 1991, shall clearly identify on any container of the system and on the system the month or week and year the system was manufactured and its conformance with the requirements of subsection (b). The display of the UL logo or listing mark, and compliance with the date marking requirements of UL—325, on both the container and the system, shall satisfy the requirements of this subsection.

(e) Notification.—Effective on and after July 1, 1991, all manufacturers of automatic residential garage door openers shall, in consultation with the Consumer Product Safety Commission, notify the public of the potential for entrapment by garage doors equipped with automatic garage door openers and advise the public to test their openers for the entrapment protection feature or device required by subsection (b).

(f) Preemption.—In applying section 26(a) of the Consumer Product Safety Act (15 U.S.C. 2075) with respect to the consumer product safety rule of the Consumer Product Safety Commission under subsection (a), only those provisions of laws of States or political subdivisions which relate to the labeling of automatic residential garage door openers and those provisions which do not provide at least the equivalent degree of protection from the risk of injury associated with automatic residential garage door openers as the consumer product safety rule provides shall be subject to such section.

(g) Regulations.—Section 553 of title 5, United States Code, shall apply with respect to the issuance of any regulations by the Consumer Product Safety Commission to implement the

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requirements of this section and sections 7 and 9 of the Consumer Product Safety Act do not apply to such issuance. Any additional or revised requirement issued by the Commission shall provide an adequate degree of protection to the public.

(h) Construction.—Nothing in this section shall affect or modify in any way the obligations or liabilities of any person under the common law or any Federal or State law.

**BICYCLE HELMETS**

{Not technically part of the Consumer Product Safety Act}

(a) In General.—Bicycle helmets manufactured 9 months or more after the date of the enactment of this Act shall conform to—
(1) any interim standard described under subsection (b), pending the establishment of a final standard pursuant to subsection (c); and
(2) the final standard, once it has been established under subsection (c).

(b) Interim Standards.—The interim standards are as follows:
(1) The American National Standards Institute standard designated as “Z90.4-1984”.
(2) The Snell Memorial Foundation standard designated as “B-90”.
(3) The American Society for Testing and Materials (ASTM) standard designated as “F 1447”.
(4) Any other standard that the Commission determines is appropriate. [16 CFR 1203]

(c) Final Standard.—Not later than 60 days after the date of the enactment of this Act, the Commission shall begin a proceeding under section 553 of title 5, United States Code, to—
(1) review the requirements of the interim standards set forth in subsection (a) and establish a final standard based on such requirements;
(2) include in the final standard a provision to protect against the risk of helmets coming off the heads of bicycle riders;
(3) include in the final standard provisions that address the risk of injury to children; and
(4) include additional provisions as appropriate. Sections 7, 9, and 30(d) of the Consumer Product Safety Act (15 U.S.C. 2056, 2058, 2079(d)) shall not apply to the proceeding under this subsection and section 11 of such Act (15 U.S.C. 2060) shall not apply with respect to any standard issued under such
proceeding. The final standard shall take effect 1 year from the
date it is issued.

(d) Failure To Meet Standards.—
(1) Failure to meet interim standard.—Until the final standard takes
effect, a bicycle helmet that does not conform to an interim
standard as required under subsection (a)(1) shall be
considered in violation of a consumer product safety standard
promulgated under the Consumer Product Safety Act.
(2) Status of final standard.—The final standard developed under
subsection (c) shall be considered a consumer product safety
standard promulgated under the Consumer Product Safety Act.

COMMISSION RESPONSIBILITY—PETITION
FOR CONSUMER PRODUCT SAFETY RULE

SEC. 10. {Repealed}

JUDICIAL REVIEW OF CONSUMER PRODUCT SAFETY RULES


(a) Not later than 60 days after a consumer product safety
rule is promulgated by the Commission, any person adversely
affected by such rule, or any consumer or consumer
organization, may file a petition with the United States court of
appeals for the District of Columbia or for the circuit in which
such person, consumer, or organization resides or has his
principal place of business for judicial review of such rule.
Copies of the petition shall be forthwith transmitted by the clerk
of the court to the Commission or other officer designated by it
for that purpose and to the Attorney General. The record of the
proceedings on which the Commission based its rule shall be
filed in the court as provided for in section 2112 of title 28,
United States Code. For purposes of this section, the term
“record” means such consumer product safety rule; any notice
or proposal published pursuant to section 7, 8, or 9; the
transcript required by section 9(d)(2) of any oral presentation;
any written submission of interested parties; and any other
information which the Commission considers relevant to such
rule.

(b) If the petitioner applies to the court for leave to adduce
additional data, views, or arguments and shows to the
satisfaction of the court that such additional data, views, or
arguments are material and that there were reasonable grounds
for the petitioner’s failure to adduce such data, views, or
arguments in the proceeding before the Commission, the court
may order the Commission to provide additional opportunity for
the oral presentation of data, views, or arguments and for

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written submissions. The Commission may modify its findings, or make new findings by reason of the additional data, views, or arguments so taken and shall file such modified or new findings, and its recommendation, if any, for the modification or setting aside of its original rule, with the return of such additional data, views, or arguments.

(c) Upon the filing of the petition under subsection (a) of this section the court shall have jurisdiction to review the consumer product safety rule in accordance with chapter 7 of title 5, United States Code, and to grant appropriate relief, including interim relief, as provided in such chapter. A court may in the interest of justice include in such relief an award of the costs of suit, including reasonable attorneys’ fees (determined in accordance with section (f))) and reasonable expert witnesses’ fees. Attorneys’ fees may be awarded against the United States (or any agency or official of the United States) without regard to section 2412 of title 28, United States Code, or any other provision of law. The consumer product safety rule shall not be affirmed unless the Commission’s findings under sections 9(f)(1) and 9(f)(3) are supported by substantial evidence on the record taken as a whole.

(d) The judgment of the court affirming or setting aside, in whole or in part, any consumer product safety rule shall be final, subject to review by the Supreme Court of the United States upon certiorari or certification, as provided in section 1254 of title 28 of the United States Code.

(e) The remedies provided for in this section shall be in addition to and not in lieu of any other remedies provided by law.

(f) For purposes of this section and sections 23(a) and 24, a reasonable attorney’s fee is a fee (1) which is based upon (A) the actual time expended by an attorney in providing advice and other legal services in connection with representing a person in an action brought under this section, and (B) such reasonable expenses as may be incurred by the attorney in the provision of such services, and (2) which is computed at the rate prevailing for the provision of similar services with respect to actions brought in the court which is awarding such fee.

IMMINENT HAZARDS


(a) The Commission may file in a United States district court an action (1) against an imminently hazardous consumer product for seizure of such product under subsection (b)(2), or (2) against any person who is a manufacturer, distributor, or retailer of such product, or (3) against both. Such an action

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may be filed notwithstanding the existence of a consumer product safety rule applicable to such product, or the pendency of any administrative or judicial proceedings under any other provision of this Act. As used in this section, and hereinafter in this Act, the term "imminently hazardous consumer product" means a consumer product which presents imminent and unreasonable risk of death, serious illness, or severe personal injury.

(b)(1) The district court in which such action is filed shall have jurisdiction to declare such product an imminently hazardous consumer product, and (in the case of an action under subsection (a)(2)) to grant (as ancillary to such declaration or in lieu thereof) such temporary or permanent relief as may be necessary to protect the public from such risk. Such relief may include a mandatory order requiring the notification of such risk to purchasers of such product known to the defendant, public notice, the recall, the repair or the replacement of, or refund for, such product.

(2) In the case of an action under subsection (a)(1), the consumer product may be proceeded against by process of libel for the seizure and condemnation of such product in any United States district court within the jurisdiction of which such consumer product is found. Proceedings and cases instituted under the authority of the preceding sentence shall conform as nearly as possible to proceedings in rem in admiralty.

(c) Where appropriate, concurrently with the filing of such action or as soon thereafter as may be practicable, the Commission shall initiate a proceeding to promulgate a consumer product safety rule applicable to the consumer product with respect to which such action is filed.

(d)(1) An action under subsection (a)(2) of this section may be brought in the United States district court for the District of Columbia or in any judicial district in which any of the defendants is found, is an inhabitant or transacts business; and process in such an action may be served on a defendant in any other district in which such defendant resides or may be found. Subpoenas requiring attendance of witnesses in such an action may run into any other district. In determining the judicial district in which an action may be brought under this section in instances in which such action may be brought in more than one judicial district, the Commission shall take into account the convenience of the parties.

(2) Whenever proceedings under this section involving substantially similar consumer products are pending in courts in two or more judicial districts, they shall be consolidated for

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trial by order of any such court upon application reasonably 
made by any party in interest, upon notice to all other parties in 
interest.

(e) Notwithstanding any other provision of law, in any action 
under this section, the Commission may direct attorneys 
employed by it to appear and represent it.

(g) Nothing in this section shall be construed to require the 
Commission, in determining whether to bring an action against 
a consumer product or a person under this section, to prepare 
a comparison of the costs that would be incurred in complying 
with the relief that may be ordered in such action with the 
benefits to the public from such relief.

RECALL OF LEAD-LINED DRINKING WATER COOLERS

[Section 1462 of the Safe Drinking Water Act, added by 
Public Law 100-572, the Lead Contamination Control act of 1988, 
[42 U.S.C. 300j-22.]
[Not technically part of the Consumer Product Safety Act]

For purposes of the Consumer Product Safety Act, all 
drinking water coolers identified by the Administrator of EPA on 
the list under section 1463 [42 U.S.C. 300j-23] as having a lead-
lined tank shall be considered to be imminently hazardous consumer products within the meaning of section 12 of such Act (15 U.S.C. 2061). After notice and opportunity for comments, including a public hearing, the Consumer Product Safety Commission shall issue an order requiring the manufacturers and importers of such coolers to repair, replace, or recall and provide a refund for such coolers within 1 year after the enactment of the Lead Contamination Control Act of 1988. [Enacted October 31, 1988] For purposes of enforcement, such an order shall be treated as an order under section 15(d) of that Act (15 U.S.C. 2064(d)).

NEW PRODUCTS

SEC. 13. [Repealed]

PRODUCT CERTIFICATION AND LABELING


(a)(1) Every manufacturer of a product which is subject to a 
consumer product safety standard under this Act and which is 
distributed in commerce (and the private labeler of such product if it bears a private label) shall issue a certificate which shall certify that such product conforms to all applicable

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consumer product safety standards, and shall specify any standard which is applicable. Such certificate shall accompany the product or shall otherwise be furnished to any distributor or retailer to whom the product is delivered. Any certificate under this subsection shall be based on a test of each product or upon a reasonable testing program; shall state the name of the manufacturer or private labeler issuing the certificate; and shall include the date and place of manufacture.

(2) In the case of a consumer product for which there is more than one manufacturer or more than one private labeler, the Commission may by rule designate one or more of such manufacturers or one or more of such private labelers (as the case may be) as the persons who shall issue the certificate required by paragraph (1) of this subsection, and may exempt all other manufacturers of such product or all other private labelers of the product (as the case may be) from the requirement under paragraph (1) to issue a certificate with respect to such product.

(b) The Commission may by rule prescribe reasonable testing programs for consumer products which are subject to consumer product safety standards under this Act and for which a certificate is required under subsection (a). Any test or testing program on the basis of which a certificate is issued under subsection (a) may, at the option of the person required to certify the product, be conducted by an independent third party qualified to perform such tests or testing programs.

(c) The Commission may by rule require the use and prescribe the form and content of labels which contain the following information (or that portion of it specified in the rule) —

(1) The date and place of manufacture of any consumer product.

(2) A suitable identification of the manufacturer of the consumer product, unless the product bears a private label in which case it shall identify the private labeler and shall also contain a code mark which will permit the seller of such product to identify the manufacturer thereof to the purchaser upon his request.

(3) In the case of a consumer product subject to a consumer product safety rule, a certification that the product meets all applicable consumer product safety standards and a specification of the standards which are applicable.

Such labels, where practicable, may be required by the Commission to be permanently marked on or affixed to any such consumer product. The Commission may, in appropriate

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cases, permit information required under paragraphs (1) and (2) of this subsection to be coded.

NOTIFICATION AND REPAIR, REPLACEMENT, OR REFUND


(a) For purposes of this section, the term “substantial product hazard” means—

(1) a failure to comply with an applicable consumer product safety rule which creates a substantial risk of injury to the public, or

(2) a product defect which (because of the pattern of defect, the number of defective products distributed in commerce, the severity of the risk, or otherwise) creates a substantial risk of injury to the public.

(b) Every manufacturer of a consumer product distributed in commerce, and every distributor and retailer of such product, who obtains information which reasonably supports the conclusion that such product—

(1) fails to comply with an applicable consumer product safety rule or with a voluntary consumer product safety standard upon which the Commission has relied under section 9;

(2) contains a defect which could create a substantial product hazard described in subsection (a)(2); or

(3) creates an unreasonable risk of serious injury or death,

shall immediately inform the Commission of such failure to comply, of such defect, or of such risk, unless such manufacturer, distributor, or retailer has actual knowledge that the Commission has been adequately informed of such defect, failure to comply, or such risk.

(c) If the Commission determines (after affording interested persons, including consumers and consumer organizations, an opportunity for a hearing in accordance with subsection (f) of this section) that a product distributed in commerce presents a substantial product hazard and that notification is required in order to adequately protect the public from such substantial product hazard, the Commission may order the manufacturer or any distributor or retailer of the product to take any one or more of the following actions:

(1) To give public notice of the defect or failure to comply.

(2) To mail notice to each person who is a manufacturer, distributor, or retailer of such product.
(3) To mail notice to every person to whom the person required to give notice knows such product was delivered or sold.

Any such order shall specify the form and content of any notice required to be given under such order.

(d) If the Commission determines (after affording interested parties, including consumers and consumer organizations, an opportunity for a hearing in accordance with subsection (f)) that a product distributed in commerce presents a substantial product hazard and that action under this subsection is in the public interest, it may order the manufacturer or any distributor or retailer of such product to take whichever of the following actions the person to whom the order is directed elects:

(1) To bring such product into conformity with the requirements of the applicable consumer product safety rule or to repair the defect in such product.

(2) To replace such product with a like or equivalent product which complies with the applicable consumer product safety rule or which does not contain the defect.

(3) To refund the purchase price of such product (less a reasonable allowance for use, if such product has been in the possession of a consumer for one year or more (A) at the time of public notice under subsection (c), or (B) at the time the consumer receives actual notice of the defect or noncompliance, whichever first occurs).

An order under this subsection may also require the person to whom it applies to submit a plan, satisfactory to the Commission, for taking action under whichever of the preceding paragraphs of this subsection under which such person has elected to act. The Commission shall specify in the order the persons to whom refunds must be made if the person to whom the order is directed elects to take the action described in paragraph (3). If an order under this subsection is directed to more than one person, the Commission shall specify which person has the election under this subsection. An order under this subsection may prohibit the person to whom it applies from manufacturing for sale, offering for sale, distributing in commerce, or importing into the customs territory of the United States (as defined in general headnote 2 to the Tariff Schedules of the United States), [19 U.S.C. 1202] or from doing any combination of such actions, the product with respect to which the order was issued.

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(e)(1) No charge shall be made to any person (other than a manufacturer, distributor, or retailer) who avails himself of any remedy provided under an order issued under subsection (d), and the person subject to the order shall reimburse each person (other than a manufacturer, distributor, or retailer) who is entitled to such a remedy for any reasonable and foreseeable expenses incurred by such person in availing himself of such remedy.

(2) An order issued under subsection (c) or (d) with respect to a product may require any person who is a manufacturer, distributor, or retailer of the product to reimburse any other person who is a manufacturer, distributor, or retailer of such product for such other person’s expenses in connection with carrying out the order, if the Commission determines such reimbursement to be in the public interest.

(f) An order under subsection (c) or (d) may be issued only after an opportunity for a hearing in accordance with section 554 of title 5, United States Code, except that, if the Commission determines that any person who wishes to participate in such hearing is a part of a class of participants who share an identity of interest, the Commission may limit such person’s participation in such hearing to participation through a single representative designated by such class (or by the Commission if such class fails to designate such a representative). Any settlement offer which is submitted to the presiding officer at a hearing under this subsection shall be transmitted by the officer to the Commission for its consideration unless the settlement offer is clearly frivolous or duplicative of offers previously made.

(g)(1) If the Commission has initiated a proceeding under this section for the issuance of an order under subsection (d) with respect to a product which the Commission has reason to believe presents a substantial product hazard, the Commission (without regard to section 27(b)(7)), or the Attorney General may, in accordance with section 12(d)(1), apply to a district court of the United States for the issuance of a preliminary injunction to restrain the distribution in commerce of such product pending the completion of such proceeding. If such a preliminary injunction has been issued, the Commission (or the Attorney General if the preliminary injunction was issued upon an application of the Attorney General) may apply to the issuing court for extensions of such preliminary injunction.

(2) Any preliminary injunction, and any extension of a preliminary injunction, issued under this subsection with respect to a product shall be in effect for such period as the issuing court prescribes not to exceed a period which extends beyond the thirtieth day from the date of the issuance of the

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preliminary injunction (or, in the case of a preliminary injunction which has been extended, the date of its extension) or the date of the completion or termination of the proceeding under this section respecting such product, whichever date occurs first.

(3) The amount in controversy requirement of section 1331 of title 28, United States Code, does not apply with respect to the jurisdiction of a district court of the United States to issue or extend a preliminary injunction under this subsection.

(h) Nothing in this section shall be construed to require the Commission, in determining that a product distributed in commerce presents a substantial product hazard and that notification or other action under this section should be taken, to prepare a comparison of the costs that would be incurred in providing notification or taking other action under this section with the benefits from such notification or action.

SMALL PARTS INCIDENT REPORTING

[Sec. 102 of Public Law 103-267, 108 Stat. 722, June 16, 1994]

(a) Reports to Consumer Product Safety Commission—.

(1) Requirement to report.—Each manufacturer, distributor, retailer, and importer of a marble, small ball, or latex balloon, or a toy or game that contains a marble, small ball, latex balloon, or other small part, shall report to the Commission any information obtained by such manufacturer, distributor, retailer, or importer which reasonably supports the conclusion that—

(A) an incident occurred in which a child (regardless of age) choked on such a marble, small ball, or latex balloon or on a marble, small ball, latex balloon, or other small part contained in such toy or game; and

(B) as a result of that incident the child died, suffered serious injury, ceased breathing for any length of time, or was treated by a medical professional.

(2) Treatment under cpsa.—For purposes of section 19(a)(3) of the Consumer Product Safety Act (15 U.S.C. 2068(a)(3)), the requirement to report information under this subsection is deemed to be a requirement under such Act.
(3) **Effect on liability.**—A report by a manufacturer, distributor, retailer, or importer under paragraph (1) shall not be interpreted, for any purpose, as an admission of liability or of the truth of the information contained in the report.

(b) **Confidentiality Protections.**—The confidentiality protections of section 6(b) of the Consumer Product Safety Act (15 U.S.C. 2055(b)) apply to any information reported to the Commission under subsection (a) of this section. For purposes of section 6(b)(5) of such Act, information so reported shall be treated as information submitted pursuant to section 15(b) of such Act respecting a consumer product.

**INSPECTION AND RECORDKEEPING**


(a) For purposes of implementing this Act, or rules or orders prescribed under this Act, officers or employees duly designated by the Commission, upon presenting appropriate credentials and a written notice from the Commission to the owner, operator, or agent in charge, are authorized—

1. to enter, at reasonable times, (A) any factory, warehouse, or establishment in which consumer products are manufactured or held, in connection with distribution in commerce, or (B) any conveyance being used to transport consumer products in connection with distribution in commerce; and

2. to inspect, at reasonable times and in a reasonable manner such conveyance or those areas of such factory, warehouse, or establishment where such products are manufactured, held, or transported and which may relate to the safety of such products. Each such inspection shall be commenced and completed with reasonable promptness.

(b) Every person who is a manufacturer, private labeler, or distributor of a consumer product shall establish and maintain such records, make such reports, and provide such information as the Commission may, by rule, reasonably require for the purposes of implementing this Act, or to determine compliance with rules or orders prescribed under this Act. Upon request of an officer or employee duly designated by the Commission, every such manufacturer, private labeler, or distributor shall permit the inspection of appropriate books, records, and papers relevant to determining whether such manufacturer, private labeler, or distributor has acted or is acting in compliance with this Act and rules under this Act.

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CONSUMER PRODUCT SAFETY ACT

IMPORTED PRODUCTS


(a) Any consumer product offered for importation into the customs territory of the United States (as defined in general headnote 2 to the Tariff Schedules of the United States) [19 U.S.C. 1202] shall be refused admission into such customs territory if such product—

(1) fails to comply with an applicable consumer product safety rule;

(2) is not accompanied by a certificate required by section 14, or is not labeled in accordance with regulations under section 14(c);

(3) is or has been determined to be an imminently hazardous consumer product in a proceeding brought under section 12;

(4) has a product defect which constitutes a substantial product hazard (within the meaning of section 15(a)(2)); or

(5) is a product which was manufactured by a person who the Commission has informed the Secretary of the Treasury is in violation of subsection (g).

(b) The Secretary of the Treasury shall obtain without charge and deliver to the Commission, upon the latter's request, a reasonable number of samples of consumer products being offered for import. Except for those owners or consignees who are or have been afforded an opportunity for a hearing in a proceeding under section 12 with respect to an imminently hazardous product, the owner or consignee of the product shall be afforded an opportunity by the Commission for a hearing in accordance with section 554 of title 5 of the United States Code with respect to the importation of such products into the customs territory of the United States. If it appears from examination of such samples or otherwise that a product must be refused admission under the terms of subsection (a), such product shall be refused admission, unless subsection (c) of this section applies and is complied with.

(c) If it appears to the Commission that any consumer product which may be refused admission pursuant to subsection (a) of this section can be so modified that it need not (under the terms of paragraphs (1) through (4) of subsection (a)) be refused admission, the Commission may defer final determination as to the admission of such product and, in accordance with such regulations as the Commission and the Secretary of the Treasury shall jointly agree to, permit such product to be delivered from customs custody under bond for

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the purpose of permitting the owner or consignee an opportunity to so modify such product.

(d) All actions taken by an owner or consignee to modify such product under subsection (c) shall be subject to the supervision of an officer or employee of the Commission and of the Department of the Treasury. If it appears to the Commission that the product cannot be so modified or that the owner or consignee is not proceeding satisfactorily to modify such product, it shall be refused admission into the customs territory of the United States, and the Commission may direct the Secretary to demand redelivery of the product into customs custody, and to seize the product in accordance with section 22(b) if it is not so redelivered.

(e) Products refused admission into the customs territory of the United States under this section must be exported, except that upon application, the Secretary of the Treasury may permit the destruction of the product in lieu of exportation. If the owner or consignee does not export the product within a reasonable time, the Department of the Treasury may destroy the product.

(f) All expenses (including travel, per diem or subsistence, and salaries of officers or employees of the United States) in connection with the destruction provided for in this section (the amount of such expenses to be determined in accordance with regulations of the Secretary of the Treasury) and all expenses in connection with the storage, cartage, or labor with respect to any consumer product refused admission under this section, shall be paid by the owner or consignee and, in default of such payment, shall constitute a lien against any future importations made by such owner or consignee.

(g) The Commission may, by rule, condition the importation of a consumer product on the manufacturer's compliance with the inspection and recordkeeping requirements of this Act and the Commission's rules with respect to such requirements.

(h)(1) The Commission shall establish and maintain a permanent product surveillance program, in cooperation with other appropriate Federal agencies, for the purpose of carrying out the Commission's responsibilities under this Act and the other Acts administered by the Commission and preventing the entry of unsafe consumer products into the commerce of the United States.

(2) The Commission may provide to the agencies with which it is cooperating under paragraph (1) such information, data, violator lists, test results, and other support, guidance, and documents as may be necessary or helpful for such agencies to cooperate with the Commission to carry out the product surveillance program under paragraph (1).

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(3) The Commission shall periodically report to the Congress the results of the surveillance program under paragraph (1).

EXPORTS


(a) This Act shall not apply to any consumer product if (1) it can be shown that such product is manufactured, sold, or held for sale for export from the United States (or that such product was imported for export), unless (A) such consumer product is in fact distributed in commerce for use in the United States, or (B) the Commission determines that exportation of such product presents an unreasonable risk of injury to consumers within the United States, and (2) such consumer product when distributed in commerce, or any container in which it is enclosed when so distributed, bears a stamp or label stating that such consumer product is intended for export; except that this Act shall apply to any consumer product manufactured for sale, offered for sale, or sold for shipment to any installation of the United States located outside the United States.

(b) Not less than thirty days before any person exports to a foreign country any product—

(1) which is not in conformity with an applicable consumer product safety standard in effect under this Act, or

(2) which is declared to be a banned hazardous substance by a rule promulgated under section 9, such person shall file a statement with the Commission notifying the Commission of such exportation, and the Commission, upon receipt of such statement, shall promptly notify the government of such country of such exportation and the basis for such safety standard or rule. Any statement filed with the Commission under the preceding sentence shall specify the anticipated date of shipment of such product, the country and port of destination of such product, and the quantity of such product that will be exported, and shall contain such other information as the Commission may by regulation require. Upon petition filed with the Commission by any person required to file a statement under this subsection respecting an exportation, the Commission may, for good cause shown, exempt such person from the requirement of this subsection that such a statement be filed no less than thirty days before the date of the exportation, except that in no case shall the Commission permit such a statement to be filed later than the tenth day before such date.

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PROHIBITED ACTS


(a) It shall be unlawful for any person to—

(1) manufacture for sale, offer for sale, distribute in commerce, or import into the United States any consumer product which is not in conformity with an applicable consumer product safety standard under this Act;

(2) manufacture for sale, offer for sale, distribute in commerce, or import into the United States any consumer product which has been declared a banned hazardous product by a rule under this Act;

(3) fail or refuse to permit access to or copying of records, or fail or refuse to establish or maintain records, or fail or refuse to make reports or provide information, or fail or refuse to permit entry or inspection, as required under this Act or rule thereunder;

(4) fail to furnish information required by section 15(b);

(5) fail to comply with an order issued under section 15 (c) or (d) (relating to notification, to repair, replacement, and refund, and to prohibited acts);

(6) fail to furnish a certificate required by section 14 or issue a false certificate if such person in the exercise of due care has reason to know that such certificate is false or misleading in any material respect; or to fail to comply with any rule under section 14(c) (relating to labeling);

(7) fail to comply with any rule under section 9(g)(2) (relating to stockpiling);

(8) fail to comply with any rule under section 13

(9) fail to comply with any rule under section 27(e) (relating to provision of performance and technical data); and

(10) fail to file a statement with the Commission pursuant to section 18(b).

(b) Paragraphs (1) and (2) of subsection (a) of this section shall not apply to any person (1) who holds a certificate issued in accordance with section 14(a) to the effect that such consumer product conforms to all applicable consumer product

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safety rules, unless such person knows that such consumer product does not conform, or (2) who relies in good faith on the representation of the manufacturer or a distributor of such product that the product is not subject to an applicable product safety rule.

**CIVIL PENALTIES**

**SEC. 20. [15 U.S.C. 2069] penalties increased; see 64 FR 51963]**

(a)(1) Any person who knowingly violates section 19 of this Act shall be subject to a civil penalty not to exceed $5,000 for each such violation. Subject to paragraph (2), a violation of section 19(a) (1), (2), (4), (5), (6), (7), (8), (9), (10), or (11) shall constitute a separate offense with respect to each consumer product involved, except that the maximum civil penalty shall not exceed $1,250,000 for any related series of violations. A violation of section 19(a)(3) shall constitute a separate violation with respect to each failure or refusal to allow or perform an act required thereby; and, if such violation is a continuing one, each day of such violations shall constitute a separate offense, except that the maximum civil penalty shall not exceed $1,250,000 for any related series of violations.

(2) The second sentence of paragraph (1) of this subsection shall not apply to violations of paragraph (1) or (2) of section 19(a)—

(A) if the person who violated such paragraphs is not the manufacturer or private labeler or a distributor of the products involved, and

(B) if such person did not have either (i) actual knowledge that his distribution or sale of the product violated such paragraphs or (ii) notice from the Commission that such distribution or sale would be a violation of such paragraphs.

(3)(A) The maximum penalty amounts authorized in paragraph (1) shall be adjusted for inflation as provided in this paragraph.

(B) Not later than December 1, 1994, and December 1 of each fifth calendar year thereafter, the Commission shall prescribe and publish in the Federal Register a schedule of maximum authorized penalties that shall apply for violations that occur after January 1 of the year immediately following such publication.

(C) The schedule of maximum authorized penalties shall be prescribed by increasing each of the amounts referred to in paragraph (1) by the cost-of-living

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adjustment for the preceding five years. Any increase determined under the preceding sentence shall be rounded to—

(i) in the case of penalties greater than $1,000 but less than or equal to $10,000, the nearest multiple of $1,000;

(ii) in the case of penalties greater than $10,000 but less than or equal to $100,000, the nearest multiple of $5,000;

(iii) in the case of penalties greater than $100,000 but less than or equal to $200,000, the nearest multiple of $10,000; and

(iv) in the case of penalties greater than $200,000, the nearest multiple of $25,000.

(D) For purposes of this subsection:

(i) The term “Consumer Price Index” means the Consumer Price Index for all-urban consumers published by the Department of Labor.

(ii) The term “cost-of-living adjustment for the preceding five years” means the percentage by which—

(I) the Consumer Price Index for the month of June of the calendar year preceding the adjustment; exceeds

(II) the Consumer Price Index for the month of June preceding the date on which the maximum authorized penalty was last adjusted.

(b) In determining the amount of any penalty to be sought upon commencing an action seeking to assess a penalty for a violation of section 19(a), the Commission shall consider the nature of the product defect, the severity of the risk of injury, the occurrence of absence of injury, the number of defective products distributed, and the appropriateness of such penalty in relation to the size of the business of the person charged.

(c) Any civil penalty under this section may be compromised by the Commission. In determining the amount of such penalty or whether it should be remitted or mitigated and in what amount, the Commission shall consider the appropriateness of such penalty to the size of the business of the person charged, the nature of the product defect, the severity of the risk of injury, the occurrence or absence of injury, and the number of defective products distributed. The amount of such penalty when finally determined, or the amount agreed on compromise, may be deducted from any sums owing by the United States to the person charged.

(d) As used in the first sentence of subsection (a)(1) of this section, the term “knowingly” means (1) the having of actual

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knowledge, or (2) the presumed having of knowledge deemed to be possessed by a reasonable man who acts in the circumstances, including knowledge obtainable upon the exercise of due care to ascertain the truth of representations.

REPORT ON CIVIL PENALTIES

[Sec. 115(d) of Public Law 101-608]
{Not technically part of the Consumer Product Safety Act}

(1) Beginning 1 year after the date of enactment of this Act, {Enacted November 16, 1990} and every year thereafter, the Consumer Product Safety Commission shall submit to the Committee on Commerce, Science, and Transportation of the Senate and the Committee on Energy and Commerce {now Committee on Commerce} of the House of Representatives the information specified in paragraph (2) of this subsection. Such information may be included in the annual report to the Congress submitted by the Commission.

(2) The Commission shall submit information with respect to the imposition of civil penalties under the statutes which it administers. The information shall include the number of civil penalties imposed, and identification of the violations that led to the imposition of such penalties, and the amount of revenue recovered from the imposition of such penalties.

CRIMINAL PENALTIES


(a) Any person who knowingly and willfully violates section 19 of this Act after having received notice of noncompliance from the Commission shall be fined not more than $50,000 or be imprisoned not more than one year, or both.

{Modified by 18 U.S.C. 3571 as follows—
Organizations:
Not more than $200,000 if the offense does not result in death.
Not more than $500,000 if the offense results in death.
Individuals:
Not more than $100,000 if the offense does not result in death.
Not more than $250,000 if the offense results in death.}

(b) Any individual director, officer, or agent of a corporation who knowingly and willfully authorizes, orders, or performs any of the acts or practices constituting in whole or in part a violation of section 19, and who has knowledge of notice of noncompliance received by the corporation from the Commission, shall be subject to penalties under this section without regard to any penalties to which that corporation may be subject under subsection (a).

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CONSUMER PRODUCT SAFETY ACT

INJUNCTIVE ENFORCEMENT AND SEIZURE


(a) The United States district courts shall have jurisdiction to take the following action:

(1) Restrain any violation of section 19.

(2) Restrain any person from manufacturing for sale, offering for sale, distributing in commerce, or importing into the United States a product in violation of an order in effect under section 15(d).

(3) Restrain any person from distributing in commerce a product which does not comply with a consumer product safety rule.

Such actions may be brought by the Commission (without regard to section 27(b)(7)(A)) or by the Attorney General in any United States district court for a district wherein any act, omission, or transaction constituting the violation occurred, or in such court for the district wherein the defendant is found or transacts business. In any action under this section process may be served on a defendant in any other district in which the defendant resides or may be found.

(b) Any consumer product—

(1) which fails to conform with an applicable consumer product safety rule, or

(2) the manufacture for sale, offering for sale, distribution in commerce, or the importation into the United States of which has been prohibited by an order in effect under section 15(d),

when introduced into or while in commerce or while held for sale after shipment in commerce shall be liable to be proceeded against on libel of information and condemned in any district court of the United States within the jurisdiction of which such consumer product is found. Proceedings in cases instituted under the authority of this subsection shall conform as nearly as possible to proceedings in rem in admiralty. Whenever such proceedings involving substantially similar consumer products are pending in courts of two or more judicial districts they shall be consolidated for trial by order of any such court upon application reasonably made by any party in interest upon notice to all other parties in interest.

SUITES FOR DAMAGES BY PERSONS INJURED


(a) Any person who shall sustain injury by reason of any knowing (including willful) violation of a consumer product

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safety rule, or any other rule or order issued by the Commission may sue any person who knowingly (including willfully) violated any such rule or order in any district court of the United States in the district in which the defendant resides or is found or has an agent, shall recover damages sustained, and may, if the court determines it to be in the interest of justice, recover the costs of suit, including reasonable attorneys’ fees (determined in accordance with section 11(f)) and reasonable expert witnesses’ fees:

Provided, That the matter in controversy exceeds the sum or value of $10,000, exclusive of interest and costs, unless such action is brought against the United States, any agency thereof, or any officer or employee thereof in his official capacity.

(b) Except when express provision is made in a statute of the United States, in any case in which the plaintiff is finally adjudged to be entitled to recover less than the sum or value of $10,000, computed without regard to any setoff or counterclaim to which the defendant may be adjudged to be entitled, and exclusive of interests and costs, the district court may deny costs to the plaintiff and, in addition, may impose costs on the plaintiff.

(c) The remedies provided for in this section shall be in addition to and not in lieu of any other remedies provided by common law or under Federal or State law.

PRIVATE ENFORCEMENT OF PRODUCT SAFETY RULES AND OF SECTION 15 ORDERS


Any interested person (including any individual or nonprofit, business, or other entity) may bring an action in any United States district court for the district in which the defendant is found or transacts business to enforce a consumer product safety rule or an order under section 15, and to obtain appropriate injunctive relief. Not less than thirty days prior to the commencement of such action, such interested person shall give notice by registered mail to the Commission, to the Attorney General, and to the person against whom such action is directed. Such notice shall state the nature of the alleged violation of any such standard or order, the relief to be requested, and the court in which the action will be brought. No separate suit shall be brought under this section if at the time the suit is brought the same alleged violation is the subject of a pending civil or criminal action by the United States under this Act. In any action under this section the court may in the

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interest of justice award the costs of suit, including reasonable attorneys’ fees (determined in accordance with section 11(f)) and reasonable expert witnesses’ fees.

EFFECT ON PRIVATE REMEDIES


(a) Compliance with consumer product safety rules or other rules or orders under this Act shall not relieve any person from liability at common law or under State statutory law to any other person.

(b) The failure of the Commission to take any action or commence a proceeding with respect to the safety of a consumer product shall not be admissible in evidence in litigation at common law or under State statutory law relating to such consumer product.

(c) Subject to sections 6(a)(2) and 6(b) but notwithstanding section 6(a)(1), (1) any accident or investigation report made under this Act by an officer or employee of the Commission shall be made available to the public in a manner which will not identify any injured person or any person treating him, without the consent of the person so identified, and (2) all reports on research projects, demonstration projects, and other related activities shall be public information.

EFFECT ON STATE STANDARDS


(a) Whenever a consumer product safety standard under this Act is in effect and applies to a risk of injury associated with a consumer product, no State or political subdivision of a State shall have any authority either to establish or to continue in effect any provision of a safety standard or regulation which prescribes any requirements as to the performance, composition, contents, design, finish, construction, packaging, or labeling of such product which are designed to deal with the same risk of injury associated with such consumer product, unless such requirements are identical to the requirements of the Federal standard.

(b) Subsection (a) of this section does not prevent the Federal Government or the government of any State or political subdivision of a State from establishing or continuing in effect a safety requirement applicable to a consumer product for its own use which requirement is designed to protect against a risk of injury associated with the product and which is not identical to the consumer product safety standard applicable to the product under this Act if the Federal, State, or political subdivision

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requirement provides a higher degree of protection from such risk of injury than the standard applicable under this Act.

(c) Upon application of a State or political subdivision of a State, the Commission may by rule, after notice and opportunity for oral presentation of views, exempt from the provisions of subsection (a) (under such conditions as it may impose in the rule) any proposed safety standard or regulation which is described in such application and which is designed to protect against a risk of injury associated with a consumer product subject to a consumer product safety standard under this Act if the State or political subdivision standard or regulation—

(1) provides a significantly higher degree of protection from such risk of injury than the consumer product safety standard under this Act, and

(2) does not unduly burden interstate commerce.

In determining the burden, if any, of a State or political subdivision standard or regulation on interstate commerce, the Commission shall consider and make appropriate (as determined by the Commission in its discretion) findings on the technological and economic feasibility of complying with such standard or regulation, the cost of complying with such standard or regulation, the geographic distribution of the consumer product to which the standard or regulation would apply, the probability of other States or political subdivisions applying for an exemption under this subsection for a similar standard or regulation, and the need for a national, uniform standard under this Act for such consumer product.

ADDITIONAL FUNCTIONS OF COMMISSION


(a) The Commission may, by one or more of its members or by such agents or agency as it may designate, conduct any hearing or other inquiry necessary or appropriate to its functions anywhere in the United States. A Commissioner who participates in such a hearing or other inquiry shall not be disqualified solely by reason of such participation from subsequently participating in a decision of the Commission in the same matter. The Commission shall publish notice of any proposed hearing in the Federal Register and shall afford a reasonable opportunity for interested persons to present relevant testimony and data.

(b) The Commission shall also have the power—

(1) to require, by special or general orders, any person to submit in writing such reports and answers to questions as the Commission may prescribe to carry

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out a specific regulatory or enforcement function of the Commission; and such submission shall be made within such reasonable period and under oath or otherwise as the Commission may determine;

(2) to administer oaths;

(3) to require by subpoena the attendance and testimony of witnesses and the production of all documentary evidence relating to the execution of its duties;

(4) in any proceeding or investigation to order testimony to be taken by deposition before any person who is designated by the Commission and has the power to administer oaths and, in such instances, to compel testimony and the production of evidence in the same manner as authorized under paragraph (3) of this subsection;

(5) to pay witnesses the same fees and mileage as are paid in like circumstances in the courts of the United States;

(6) to accept gifts and voluntary and uncompensated services, notwithstanding the provisions of section 3679 of the Revised Statutes (31 U.S.C. 665 (b)); [Now 31 U.S.C. 1342]

(7) to—

(A) initiate, prosecute, defend, or appeal (other than to the Supreme Court of the United States), through its own legal representative and in the name of the Commission, any civil action if the Commission makes a written request to the Attorney General for representation in such civil action and the Attorney General does not within the 45-day period beginning on the date such request was made notify the Commission in writing that the Attorney General will represent the Commission in such civil action, and

(B) initiate, prosecute, or appeal, through its own legal representative, with the concurrence of the Attorney General or through the Attorney General, any criminal action, for the purpose of enforcing the laws subject to its jurisdiction;

(8) to lease buildings or parts of buildings in the District of Columbia, without regard to the Act of March 3, 1877 (40 U.S.C. 34), for the use of the Commission; and

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(9) to delegate any of its functions or powers, other than the power to issue subpenas under paragraph (3), to any officer or employee of the Commission.

An order issued under paragraph (1) shall contain a complete statement of the reason the Commission requires the report or answers specified in the order to carry out a specific regulatory or enforcement function of the Commission. Such an order shall be designed to place the least burden on the person subject to the order as is practicable taking into account the purpose for which the order was issued.

(c) Any United States district court within the jurisdiction of which any inquiry is carried on, may, upon petition by the Commission (subject to subsection (b)(7)) or by the Attorney General, in case of refusal to obey a subpena or order of the Commission issued under subsection (b) of this section, issue an order requiring compliance therewith; and any failure to obey the order of the court may be punished by the court as a contempt thereof.

(d) No person shall be subject to civil liability to any person (other than the Commission or the United States) for disclosing information at the request of the Commission.

(e) The Commission may by rule require any manufacturer of consumer products to provide to the Commission such performance and technical data related to performance and safety as may be required to carry out the purposes of this Act, and to give such notification of such performance and technical data at the time of original purchase to prospective purchasers and to the first purchaser of such product for purposes other than resale, as it determines necessary to carry out the purposes of this Act.

(f) For purposes of carrying out this Act, the Commission may purchase any consumer product and it may require any manufacturer, distributor, or retailer of a consumer product to sell the product to the Commission at manufacturer's, distributor's, or retailer's cost.

(g) The Commission is authorized to enter into contracts with governmental entities, private organizations, or individuals for the conduct of activities authorized by this Act.

(h) The Commission may plan, construct, and operate a facility or facilities suitable for research, development, and testing of consumer products in order to carry out this Act.

(i)(1) Each recipient of assistance under this Act pursuant to grants or contracts entered into under other than competitive bidding procedures shall keep such records as the Commission by rule shall prescribe, including records which fully disclose the amount and disposition by such recipient of the proceeds of

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such assistance, the total cost of the project undertaken in connection with which such assistance is given or used, and the amount of that portion of the cost of the project or undertaking supplied by other sources, and such other records as will facilitate an effective audit.

(2) The Commission and the Comptroller General of the United States, or their duly authorized representatives, shall have access for the purpose of audit and examination to any books, documents, papers, and records of the recipients that are pertinent to the grants or contracts entered into under this Act under other than competitive bidding procedures.

(j) The Commission shall prepare and submit to the President and the Congress at the beginning of each regular session of Congress a comprehensive report on the administration of this Act for the preceding fiscal year. Such report shall include—

(1) a thorough appraisal, including statistical analyses, estimates, and long-term projections, of the incidence of injury and effects to the population resulting from consumer products, with a breakdown, insofar as practicable, among the various sources of such injury;

(2) a list of consumer product safety rules prescribed or in effect during such year;

(3) an evaluation of the degree of observance of consumer product safety rules, including a list of enforcement actions, court decisions, and compromises of alleged violations, by location and company name;

(4) a summary of outstanding problems confronting the administration of this Act in order of priority;

(5) an analysis and evaluation of public and private consumer product safety research activities;

(6) a list, with a brief statement of the issues, of completed or pending judicial actions under this Act;

(7) the extent to which technical information was disseminated to the scientific and commercial communities and consumer information was made available to the public;

(8) the extent of cooperation between Commission officials and representatives of industry and other interested parties in the implementation of this Act, including a log or summary of meetings held between Commission officials and representatives of industry and other interested parties;

(9) an appraisal of significant actions of State and local governments relating to the responsibilities of the Commission;

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(10) with respect to voluntary consumer product safety standards for which the Commission has participated in the development through monitoring or offering of assistance and with respect to voluntary consumer product safety standards relating to risks of injury that are the subject of regulatory action by the Commission, a description of—

(A) the number of such standards adopted;
(B) the nature and number of the products which are the subject of such standards;
(C) the effectiveness of such standards in reducing potential harm from consumer products;
(D) the degree to which staff members of the Commission participate in the development of such standards;
(E) the amount of resources of the Commission devoted to encouraging development of such standards; and
(F) such other information as the Commission determine appropriate or necessary to inform the Congress on the current status of the voluntary consumer product safety standard program; and

(11) such recommendations for additional legislation as the Commission deems necessary to carry out the purposes of this Act.

{This reporting requirement ceased to be effective with respect to Congress on December 21, 1999 per Pub. L. 104-66, § 3003.}

(k)(1) Whenever the Commission submits any budget estimate or request to the President or the Office of Management and Budget, it shall concurrently transmit a copy of that estimate or request to the Congress. {This reporting requirement ceased to be effective with respect to Congress on December 21, 1999 per Pub. L. 104-66, § 3003.}

(2) Whenever the Commission submits any legislative recommendations, or testimony, or comments on legislation to the President or the Office of Management and Budget, it shall concurrently transmit a copy thereof to the Congress.

{This reporting requirement ceased to be effective on December 21, 1999 per Pub. L. 104-66, § 3003.} No officer or agency of the United States shall have any authority to require the Commission to submit its legislative recommendations, or testimony, or comments on legislation, to any officer or agency of the United States for approval, comments, or review, prior to the submission of such recommendations, testimony, or comments to the Congress.

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(a) The Commission shall appoint Chronic Hazard Advisory Panels (hereinafter referred to as the Panel or Panels) to advise the Commission in accordance with the provisions of section 31(b) respecting the chronic hazards of cancer, birth defects, and gene mutations associated with consumer products.

(b) Each Panel shall consist of 7 members appointed by the Commission from a list of nominees who shall be nominated by the President of the National Academy of Sciences from scientists—

(1) who are not officers or employees of the United States (other than employees of the National Institutes of Health, the National Toxicology Program, or the National Center for Toxicological Research), and who do not receive compensation from or have any substantial financial interest in any manufacturer, distributor, or retailer of a consumer product; and

(2) who have demonstrated the ability to critically assess chronic hazards and risks to human health presented by the exposure of humans to toxic substances or as demonstrated by the exposure of animals to such substances.

The President of the National Academy of Sciences shall nominate for each Panel a number of individuals equal to three times the number of members to be appointed to the Panel.

(c) The Chairman and Vice Chairman of the Panel shall be elected from among the members and shall serve for the duration of the Panel.

(d) Decisions of the Panel shall be made by a majority of the Panel.

(e) The Commission shall provide each Panel with such administrative support services as it may require to carry out its duties under section 31.

(f) A member of a Panel appointed under subsection (a) shall be paid at a rate not to exceed the daily equivalent of the annual rate of basic pay in effect for grade GS-18 of the General Schedule for each day (including traveltime) during which the member is engaged in the actual performance of the duties of the Panel.

(g) Each Panel shall request information and disclose information to the public, as provided in subsection (h), only through the Commission.

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(h)(1) Notwithstanding any statutory restriction on the authority of agencies and departments of the Federal Government to share information, such agencies and departments shall provide the Panel with such information and data as each Panel, through the Commission, may request to carry out its duties under section 31. Each Panel may request information, through the Commission, from States, industry and other private sources as it may require to carry out its responsibilities.

(2) Section 6 shall apply to the disclosure of information by the Panel but shall not apply to the disclosure of information to the Panel.

§See section 31(b) for additional provisions.}

COOPERATION WITH STATES AND WITH OTHER FEDERAL AGENCIES


(a) The Commission shall establish a program to promote Federal-State cooperation for the purposes of carrying out this Act. In implementing such program the Commission may—

   (1) accept from any State or local authorities engaged in activities relating to health, safety, or consumer protection assistance in such functions as injury data collection, investigation, and educational programs, as well as other assistance in the administration and enforcement of this Act which such States or localities may be able and willing to provide and, if so agreed, may pay in advance or otherwise for the reasonable cost of such assistance, and

   (2) commission any qualified officer or employee of any State or local agency as an officer of the Commission for the purpose of conducting examinations, investigations, and inspections.

(b) In determining whether such proposed State and local programs are appropriate in implementing the purposes of this Act, the Commission shall give favorable consideration to programs which establish separate State and local agencies to consolidate functions relating to product safety and other consumer protection activities.

(c) The Commission may obtain from any Federal department or agency such statistics, data, program reports, and other materials as it may deem necessary to carry out its functions under this Act. Each such department or agency may cooperate with the Commission and, to the extent permitted by law, furnish such materials to it. The Commission and the heads of other departments and agencies engaged in administering programs related to product safety shall, to the

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maximum extent practicable, cooperate and consult in order to
insure fully coordinated efforts.

(d) The Commission shall, to the maximum extent
practicable, utilize the resources and facilities of the National
Bureau of Standards \{now the National Institute of Standards and
Technology\} on a reimbursable basis, to perform research and
analyses related to risks of injury associated with consumer
products (including fire and flammability risks), to develop test
methods, to conduct studies and investigations, and to provide
technical advice and assistance in connection with the
functions of the Commission.

(e) The Commission may provide to another Federal agency
or a State or local agency or authority engaged in activities
relating to health, safety, or consumer protection, copies of any
accident or investigation report made under this Act by any
officer, employee, or agent of the Commission only if (1)
information which under section 6(a)(2) is to be considered
confidential is not included in any copy of such report which is
provided under this subsection; and (2) each Federal agency
and State and local agency and authority which is to receive
under this subsection a copy of such report provides
assurances satisfactory to the Commission that the identity of
any injured person and any person who treated an injured
person will not, without the consent of the person identified, be
included in—

(A) any copy of any such report, or
(B) any information contained in any such report,

which the agency or authority makes available to any member
of the public. No Federal agency or State or local agency or
authority may disclose to the public any information contained
in a report received by the agency or authority under this
subsection unless with respect to such information the
Commission has complied with the applicable requirements of
section 6(b).

TRANSFERS OF FUNCTIONS


(a) The functions of the Secretary of Health, Education, and
1261 et seq.) and the Poison Prevention Packaging Act of 1970
are transferred to the Commission. The functions of the
Secretary of Health, Education, and Welfare under the Federal
Food, Drug, and Cosmetic Act (15 U.S.C. 301 et seq.), to the
extent such functions relate to the administration and
enforcement of the Poison Prevention Packaging Act of 1970,
are transferred to the Commission.

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(b) The functions of the Secretary of Health, Education, and Welfare, the Secretary of Commerce, and the Federal Trade Commission under the Flammable Fabrics Act (15 U.S.C. 1191 et seq.) are transferred to the Commission. The functions of the Federal Trade Commission under the Federal Trade Commission Act, to the extent such functions relate to the administration and enforcement of the Flammable Fabrics Act, are transferred to the Commission.


(d) A risk of injury which is associated with a consumer product and which could be eliminated or reduced to a sufficient extent by action under the Federal Hazardous Substances Act, the Poison Prevention Packaging Act of 1970, or the Flammable Fabrics Act may be regulated under this Act only if the Commission by rule finds that it is in the public interest to regulate such risk of injury under this Act. Such a rule shall identify the risk of injury proposed to be regulated under this Act and shall be promulgated in accordance with section 553 of title 5, United States Code; except that the period to be provided by the Commission pursuant to subsection (c) of such section for the submission of data, views, and arguments respecting the rule shall not exceed thirty days from the date of publication pursuant to subsection (b) of such section of a notice respecting the rule.

(e)(1)(A) All personnel, property, records, obligations, and commitments which are used primarily with respect to any function transferred under the provisions of subsections (a), (b) and (c) of this section shall be transferred to the Commission, except those associated with fire and flammability research in the National Bureau of Standards. [now the National Institute of Standards and Technology] The transfer of personnel pursuant to this paragraph shall be without reduction in classification or compensation for one year after such transfer, except that the Chairman of the Commission shall have full authority to assign personnel during such one-year period in order to efficiently carry out functions transferred to the Commission under this section.

(B) Any commissioned officer of the Public Health Service who upon the day before the effective date of this section, is serving as such officer primarily in the performance of functions transferred by this Act to the Commission, may, if such officer so elects, acquire competitive status and be transferred to a competitive position in the Commission subject to subparagraph (A) of this paragraph, under the terms

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prescribed in paragraphs (3) through (8)(A) of section 15(b) of the Clean Air Amendments of 1970 (84 Stat. 1676; 42 U.S.C. 215 nt).

(2) All orders, determinations, rules, regulations, permits, contracts, certificates, licenses, and privileges (A) which have been issued, made, granted, or allowed to become effective in the exercise of functions which are transferred under this section by any department or agency, any functions of which are transferred by this section, and (B) which are in effect at the time this section takes effect, shall continue in effect according to their terms until modified, terminated, superseded, set aside, or repealed by the Commission, by any court of competent jurisdiction, or by operation of law.

(3) The provisions of this section shall not affect any proceedings pending at the time this section takes effect before any department or agency, functions of which are transferred by this section; except that such proceedings, to the extent that they relate to functions so transferred, shall be continued before the Commission. Orders shall be issued in such proceedings, appeals shall be taken therefrom, and payments shall be made pursuant to such orders, as if this section had not been enacted; and orders issued in any such proceedings shall continue in effect until modified, terminated, superseded, or repealed by the Commission, by a court of competent jurisdiction, or by operation of law.

(4) The provisions of this section shall not affect suits commenced prior to the date this section takes effect and in all such suits proceedings shall be had, appeals taken, and judgments rendered, in the same manner and effect as if this section had not been enacted; except that if before the date on which this section takes effect, any department or agency (or officer thereof in his official capacity) is a party to a suit involving functions transferred to the Commission, then such suit shall be continued by the Commission. No cause of action, and no suit, action, or other proceeding, by or against any department or agency (or officer thereof in his official capacity) functions of which are transferred by this section, shall abate by reason of the enactment of this section. Causes of actions, suits, actions, or other proceedings may be asserted by or against the United States or the Commission as may be appropriate and, in any litigation pending when this section takes effect, the court may at any time, on its own motion or that of any party, enter an order which will give effect to the provisions of this paragraph.

(f) For purposes of this section, (1) the term “function” includes power and duty, and (2) the transfer of a function, under any provision of law, of an agency or the head of a

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department shall also be a transfer of all functions under such law which are exercised by any office or officer of such agency or department.

LIMITATION ON JURISDICTION


(a) The Commission shall have no authority under this Act to regulate any risk of injury associated with a consumer product if such risk could be eliminated or reduced to a sufficient extent by actions taken under the Occupational Safety and Health Act of 1970; [29 U.S.C. 651 et seq] the Atomic Energy Act of 1954; or the Clean Air Act. [42 u.s.c. 7401 et seq] The Commission shall have no authority under this Act to regulate any risk of injury associated with electronic product radiation emitted from an electronic product (as such terms are defined by sections 355 (1) and (2) of the Public Health Service Act) [now 21 U.S.C. 360hh] if such risk of injury may be subjected to regulation under subpart 3 of part F of title III of the Public Health Service Act. [now 21 U.S.C. 360kk]

(b)(1) The Commission may not issue—

(A) an advance notice of proposed rulemaking for a consumer product safety rule,

(B) a notice of proposed rulemaking for a rule under section 27(e), or

(C) an advance notice of proposed rulemaking for regulations under section 2(q)(1) of the Federal Hazardous Substances Act,

relating to a risk of cancer, birth defects, or gene mutations from a consumer product unless a Chronic Hazard Advisory Panel, established under section 28, has, in accordance with paragraph (2), submitted a report to the Commission with respect to whether a substance contained in such product is a carcinogen, mutagen, or teratogen.

(2)(A) Before the Commission issues an advance notice of proposed rulemaking for—

(i) a consumer product safety rule,

(ii) a rule under section 27(e), or

(iii) a regulation under section 2(q)(1) of the Federal Hazardous Substances Act,

relating to a risk of cancer, birth defects, or gene mutations from a consumer product, the Commission shall request the Panel to review the scientific data and other relevant information relating to such risk to determine if any substance in the product is a carcinogen, mutagen, or a teratogen and to report its determination to the Commission.

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(B) When the Commission appoints a Panel, the Panel shall convene within 30 days after the date the final appointment is made to the Panel. The Panel shall report its determination to the Commission not later than 120 days after the date the Panel is convened or, if the Panel requests additional time, within a time period specified by the Commission. If the determination reported to the Commission states that a substance in a product is a carcinogen, mutagen, or a teratogen, the Panel shall include in its report an estimate, if such an estimate is feasible, of the probable harm to human health that will result from exposure to the substance.

(C) A Panel appointed under section 28 shall terminate when it has submitted its report unless the Commission extends the existence of the Panel.

(D) The Federal Advisory Committee Act shall not apply with respect to any Panel established under this section.

(C) Each Panel’s report shall contain a complete statement of the basis for the Panel’s determination. The Commission shall consider the report of the Panel and incorporate such report into the advance notice of proposed rulemaking and final rule.

AUTHORIZATION OF APPROPRIATIONS

SEC. 32. [15 U.S.C. 2081]

(a) There are authorized to be appropriated for the purposes of carrying out the provisions of this Act (other than the provisions of section 27(h) which authorize the planning and construction of research, development, and testing facilities) and for the purpose of carrying out the functions, powers, and duties transferred to the Commission under section 30, not to exceed—

(1) $42,000,000 for the fiscal year 1991, and
(2) $45,000,000 for the fiscal year 1992.

For payment of accumulated and accrued leave under section 5551 of title 5, United States Code, severance pay under section 5595 under such title, and any other expense related to a reduction in force in the Commission, there are authorized to be appropriated such sums as may be necessary.

(b)(1) There are authorized to be appropriated such sums as may be necessary for the planning and construction of research, development and testing facilities described in section 27(h); except that no appropriation shall be made for any such planning or construction involving an expenditure in excess of $100,000 if such planning or construction has not been approved by resolutions adopted in substantially the same form.

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by the Committee on Committee on Energy and Commerce of the House of Representatives, and by the Committee on Commerce, Science, and Transportation of the Senate. For the purpose of securing consideration of such approval the Commission shall transmit to Congress a prospectus of the proposed facility including (but not limited to)—

(A) a brief description of the facility to be planned or constructed;  
(B) the location of the facility, and an estimate of the maximum cost of the facility;  
(C) a statement of those agencies, private and public, which will use such facility, together with the contribution to be made by each such agency toward the cost of such facility; and  
(D) a statement of justification of the need for such facility.  

This reporting requirement ceased to be effective on December 21, 1999 per Pub. L. 104-66, § 3003.}

(2) The estimated maximum cost of any facility approved under this subsection as set forth in the prospectus may be increased by the amount equal to the percentage increase, if any, as determined by the Commission, in construction costs, from the date of the transmittal of such prospectus to Congress, but in no event shall the increase authorized by this paragraph exceed 10 per centum of such estimated maximum cost.

(c) No funds appropriated under subsection (a) may be used to pay any claim described in section 4(i) whether pursuant to a judgment of a court or under any award, compromise, or settlement of such claim made under section 2672 of title 28, United States Code, or under any other provision of law.

SEPARABILITY


If any provision of this Act, or the application of such provision to any person or circumstance, shall be held invalid, the remainder of this Act, or the application of such provisions to persons or circumstances other than those as to which it is held invalid, shall not be affected thereby.

EFFECTIVE DATE

SEC. 34. [15 U.S.C. 2051n]

This Act shall take effect on the sixtieth day following the date of its enactment, except—

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(1) sections 4 and 32 shall take effect on the date of enactment of this Act, and
(2) section 30 shall take effect on the later of (A) 150 days after the date of enactment of this Act, or (B) the date on which at least three members of the Commission first take office.  

INTERIM CELLULOSE INSULATION SAFETY STANDARD


(a)(1) Subject to the provisions of paragraph (2), on and after the last day of the 60-day period beginning on the effective date of this section, the requirements for flame resistance and corrosiveness set forth in the General Services Administration’s specification for cellulose insulation, HH-I-515C (as such specification was in effect on February 1, 1978), shall be deemed to be an interim consumer product safety standard which shall have all the authority and effect of any other consumer product safety standard promulgated by the Commission under this Act. During the 45-day period beginning on the effective date of this section, the Commission may make, and shall publish in the Federal Register, such technical, nonsubstantive changes in such requirements as it deems appropriate to make such requirements suitable for promulgation as a consumer product safety standard. At the end of the 60-day period specified in the first sentence of this paragraph, the Commission shall publish in the Federal Register such interim consumer product safety standard, as altered by the Commission under this paragraph.

(2) The interim consumer product safety standard established in paragraph (1) shall provide that any cellulose insulation which is produced or distributed for sale or use as a consumer product shall have a flame spread rating of 0 to 25, as such rating is set forth in the General Services Administration’s specification for cellulose insulation, HH-I-515C.

(3) During the period for which the interim consumer product safety standard established in subsection (a) is in effect, in addition to complying with any labeling requirement established by the Commission under this Act, each manufacturer or private labeler of cellulose insulation shall include the following statement on any container of such cellulose insulation: “ATTENTION: This material meets the applicable minimum Federal flammability standard. This standard is based upon laboratory tests only, which do not represent actual conditions which may occur in the home”. Such statement shall be located in a conspicuous place on such

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(b) Judicial review of the interim consumer product safety standard established in subsection (a), as such standard is in effect on and after the last day of the 60-day period specified in such subsection, shall be limited solely to the issue of whether any changes made by the Commission under paragraph (1) are technical nonsubstantive changes. For purposes of such review, any change made by the Commission under paragraph (1) which requires that any test to determine the flame spread rating of cellulose insulation shall include a correction for variations in test results caused by equipment used in the test shall be considered a technical, nonsubstantive change.

(c)(1)(A) Any interim consumer product safety standard established pursuant to this section shall be enforced in the same manner as any other consumer product safety standard until such time as there is in effect a final consumer product safety standard promulgated by the Commission, as provided in subparagraph (B), or until such time as it is revoked by the Commission under section 9(e). A violation of the interim consumer product safety standard shall be deemed to be a violation of a consumer product safety standard promulgated by the Commission under section 9.

(B) If the Commission determines that the interim consumer product safety standard does not adequately protect the public from the unreasonable risk of injury associated with flammable or corrosive cellulose insulation, it shall promulgate a final consumer product safety standard to protect against such risk. Such final standard shall be promulgated pursuant to section 553 of title 5, United States Code, except that the Commission shall give interested persons an opportunity for the oral presentation of data, views, or arguments, in addition to an opportunity to make written submissions. A transcript shall be kept of any oral presentation. The provisions of section 9(b), (c), and (d) shall apply to any proceeding to promulgate such final standard. In any judicial review of such final standard under section 11, the court shall not require any demonstration that each particular finding made by the Commission under section 9(c) is supported by substantial evidence. The court shall affirm the action of the Commission unless the court determines that such action is not supported by substantial evidence on the record taken as a whole.

(2)(A) Until there is in effect such a final consumer product safety standard, the Commission shall incorporate into the interim consumer product safety standard, in accordance with the provisions of this paragraph, each revision superseding the

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requirements for flame resistance and corrosiveness referred to in subsection (a) and promulgated by the General Services Administration.

(B) At least 45 days before any revision superseding such requirements is to become effective, the Administrator of the General Services Administration shall notify the Commission of such revision. In the case of any such revision which becomes effective during the period beginning on February 1, 1978, and ending on the effective date of this section, such notice from the Administrator of the General Services Administration shall be deemed to have been made on the effective date of this section.

(C)(i) No later than 45 days after receiving any notice under subparagraph (B), the Commission shall publish the revision, including such changes in the revision as it considers appropriate to make the revision suitable for promulgation as an amendment to the interim consumer product safety standard, in the Federal Register as a proposed amendment to the interim consumer product safety standard.

(ii) The Commission may extend the 45-day period specified in clause (i) for an additional period of not more than 150 days if the Commission determines that such extension is necessary to study the technical and scientific basis for the revision involved, or to study the safety and economic consequences of such revision.

(D)(i) Additional extensions of the 45-day period specified in subparagraph (C)(i) may be taken by the Commission if—

(I) the Commission makes the determination required in subparagraph (C)(ii) with respect to each such extension; and

(II) in the case of further extensions proposed by the Commission after an initial extension under this clause, such further extensions have not been disapproved under clause (iv).

(ii) Any extension made by the Commission under this subparagraph shall be for a period of not more than 45 days.

(iii) Prior notice of each extension made by the Commission under this subparagraph, together with a statement of the reasons for such extension and an estimate of the length of time required by the Commission to complete its action upon the revision involved, shall be published in the Federal Register and shall be submitted to the Committee on Commerce, Science, and Transportation of the Senate and the Committee on Interstate and Foreign Commerce of the House of Representatives. {This reporting requirement ceased to be effective with respect to Congress on December 21, 1999 per Pub. L. 104-66, § 3003.}
(iv) In any case in which the Commission takes an initial 45-day extension under clause (i), the Commission may not take any further extensions under clause (i) if each committee referred to in clause (iii) disapproves by committee resolution any such further extensions before the end of the 15-day period following notice of such initial extension made by the Commission in accordance with clause (iii).

(E) The Commission shall give interested persons an opportunity to comment upon any proposed amendment to the interim consumer product safety standard during the 30-day period following any publication by the Commission under subparagraph (C).

(F) No later than 90 days after the end of the period specified in subparagraph (E), the Commission shall promulgate the amendment to the interim consumer product safety standard unless the Commission determines, after consultation with the Secretary of Energy, that—

(i) such amendment is not necessary for the protection of consumers from the unreasonable risk of injury associated with flammable or corrosive cellulose insulation; or

(ii) implementation of such amendment will create an undue burden upon persons who are subject to the interim consumer product safety standard.

(G) The provisions of section 11 shall not apply to any judicial review of any amendment to the interim product safety standard promulgated under this paragraph.

(d) Any Federal department, agency, or instrumentality, or any Federal independent regulatory agency, which obtains information which reasonably indicates that cellulose insulation is being manufactured or distributed in violation of this Act shall immediately inform the Commission of such information.

(e)(1) The Commission, no later than 45 days after the effective date of this section, shall submit a report to the Committee on Commerce, Science, and Transportation of the Senate and to the Committee on Interstate and Foreign Commerce of the House of Representatives which shall contain a detailed statement of the manner in which the Commission intends to carry out the enforcement of this section.

(2)(A) The Commission, no later than 6 months after the date upon which the report required in paragraph (1) is due (and no later than the end of each 6-month period thereafter), shall submit a report to each committee referred to in
paragraph (1) which shall describe the enforcement activities of the Commission with respect to this section during the most recent 6-month period.

(B) The first report which the Commission submits under subparagraph (A) shall include the results of tests of cellulose insulation manufactured by at least 25 manufacturers which the Commission shall conduct to determine whether such cellulose insulation complies with the interim consumer product safety standard. The second such report shall include the results of such tests with respect to 50 manufacturers who were not included in testing conducted by the Commission for inclusion in the first report.

(f)(1) The Commission shall have the authority to require that any person required to comply with the certification requirements of section 14 with respect to the manufacture of cellulose insulation shall provide for the performance of any test or testing program required for such certification through the use of an independent third party qualified to perform such test or testing program. The Commission may impose such requirement whether or not the Commission has established a testing program for cellulose insulation under section 14(b).

(2) The Commission, upon petition by a manufacturer, may waive the requirements of paragraph (1) with respect to such manufacturer if the Commission determines that the use of an independent third party is not necessary in order for such manufacturer to comply with the certification requirements of section 14.

(3) The Commission may prescribe such rules as it considers necessary to carry out the provisions of this subsection.

(g) There are authorized to be appropriated, for each of the fiscal years 1978, 1979, 1980, and 1981, such sums as may be necessary to carry out the provisions of this section.

CONGRESSIONAL VETO OF CONSUMER PRODUCT SAFETY RULES


(a) The Commission shall transmit to the Secretary of the Senate and the Clerk of the House of Representatives a copy of any consumer product safety rule promulgated by the Commission under section 9.

(b) Any rule specified in subsection (a) shall not take effect if—

(1) within 90 calendar days of continuous session of the Congress which occur after the date of the promulgation of such rule, both House of the Congress

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adopt a concurrent resolution, the matter after the resolving clause of which is as follow (with the blank spaces appropriately filled): “That the Congress disapproves the consumer product safety rule which was promulgated by the Consumer Product Safety Commission with respect to and which was transmitted to the Congress on and disapproves the rule for the following reasons: ”; or

(2) within the 60 calendar days of continuous session of the Congress which occur after the date of the promulgation of such rule, one House of the Congress adopts such concurrent resolution and transmits such resolution to the other House and such resolution is not disapproved by such other House within the 30 calendar days of continuous session of the Congress which occur after the date of such transmittal.

(c) Congressional inaction on, or rejection of, a concurrent resolution of disapproval under this section shall not be construed as an expression of approval of the rule involved, and shall not be construed to create any presumption of validity with respect to such rule.

(d) For purposes of this section—

(1) continuity of session is broken only by an adjournment of the Congress sine die; and

(2) the days on which either House is not in session because of an adjournment of more than 3 days to a day certain are excluded in the computation of the periods of continuous session of the Congress specified in subsection (b).

INFORMATION REPORTING


(a) If a particular model of a consumer product is the subject of at least 3 civil actions that have been filed in Federal or State court for death or grievous bodily injury which in each of the 24-month periods defined in subsection (b) result in either a final settlement involving the manufacturer or a court judgment in favor of the plaintiff, the manufacturer of such product shall, in accordance with subsection (c), report to the Commission each such civil action within 30 days after the final settlement or court judgment in the third of such civil actions, and, within 30 days after any subsequent settlement or judgment in that 24-month period, any other such action.
(b) The 24-month periods referred to in subsection (a) are the 24-month period commencing on January 1, 1991, and subsequent 24-month periods beginning on January 1 of the calendar year that is two years following the beginning of the previous 24-month period.

(c)(1) The information required by subsection (a) to be reported to the Commission, with respect to each civil action described in subsection (a), shall include and in addition to any voluntary information provided under paragraph (2) shall be limited to the following:

(A) The name and address of the manufacturer.
(B) The model and model number or designation of the consumer product subject to the civil action.
(C) A statement as to whether the civil action alleged death or grievous bodily injury, and in the case of an allegation of grievous bodily injury, a statement of the category of such injury.
(D) A statement as to whether the civil action resulted in a final settlement or a judgment in favor of the plaintiff.
(E) In the case of a judgment in favor of the plaintiff, the number assigned, the name of the civil action, the number assigned the civil action, and the court in which the civil action was filed.

(2) A manufacturer furnishing the report required by paragraph (1) may include (A) a statement as to whether any judgment in favor of the plaintiff is under appeal or is expected to be appealed or (B) any other information which the manufacturer chooses to provide. A manufacturer reporting to the Commission under subsection (a) need not admit or may specifically deny that the information it submits reasonably supports the conclusion that its consumer product caused a death or grievous bodily injury.

(3) No statement of the amount paid by the manufacturer in a final settlement shall be required a part of the report furnished under subsection (a), nor shall such a statement of settlement amount be required under any other section of this Act.

(d) The reporting of a civil action described in subsection (a) by a manufacturer shall not constitute an admission of—

(1) an unreasonable risk of injury,
(2) a defect in the consumer product which was the subject of such action,
(3) a substantial product hazard,
(4) an imminent hazard,
(5) any other admission of liability under any statute or under any common law.

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(e) For purposes of this section:

(1) A grievous bodily injury includes any of the following categories of injury: mutilation, amputation, dismemberment, disfigurement, loss of important bodily functions, debilitating internal disorder, severe burn, severe electric shock, and injuries likely to require extended hospitalization.

(2) For purposes of this section, a particular model of a consumer product is one that is distinctive in functional design, construction, warnings or instructions related to safety, function, user population, or other characteristics which could affect the product’s safety related performance.