Consumer Product Safety Commission

This proceeding begins and ends in accordance with the applicable regulations or procedures of the administrative body before which the proceeding is heard.

(7) A proceeding to obtain a retraction from the Commission pursuant to subpart F of these rules. This proceeding begins with the filing with the Secretary of the Commission of a request for retraction and ends when the request is denied or, if granted, when the information is retracted.

In the course of or concerning. The phrase “in the course of or concerning” shall have the same meaning as set forth in either §1101.44 (c) and (d) or §1101.45 (c) and (d), whichever is applicable.

Subpart F—Retraction

§ 1101.51 Commission interpretation.

(a) Statutory provisions. Section 6(b)(7) of the CPSA provides: 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.

(b) Scope. Section 6(b)(7) applies to inaccurate or misleading information only if it is adverse—i.e., if it reflects adversely either on the safety of a consumer product or on the practices of a 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.

§ 1101.52 Procedure for retraction.

(a) Initiative. The Commission may retract information under section 6(b)(7) on the initiative of the Commission, upon the request of a manufacturer, private labeler, distributor, or retailer of a consumer product, or upon the request of any other person in accordance with the procedures provided in this section.

(b) Request for retraction. Any manufacturer, private labeler, distributor or retailer of a consumer product or any other person may request a retraction if he/she believes the Commission or an individual member, employee, agent, contractor or representative of the Commission has made public disclosure of inaccurate or misleading information, which reflects adversely either on the safety of a product with which the firm deals or on the practices of the firm. The request must be in writing and addressed to the Secretary, CPSC, Washington, D.C. 20207.

(c) Content of request. A request for retraction must include the following information to the extent it is reasonably available:

(1) The information disclosed for which retraction is requested, the date on which the information was disclosed, and the type of document (e.g., letter, memorandum, news release) and any other relevant information the firm has to assist the Commission in identifying the information. A photocopy of the disclosure should accompany the request.

(2) A statement of the specific aspects of the information the firm believes are inaccurate or misleading and reflect adversely either on the safety of a consumer product with which the firm deals or on the firm’s practices.

(3) A statement of the reasons the firm believes the information is inaccurate or misleading and reflects adversely either on the safety of a consumer product with which the firm deals or on the firm’s practices.

(4) A statement of the action the firm requests the Commission to take in publishing a retraction in a manner equivalent to that in which disclosure was made.

(5) Any additional data or information the firm believes is relevant.
§ 1101.61 Generally.

(a) Generally. In addition to the requirements of section 6(b)(1), section 6(b)(5) of the CPSA imposes further limitations on the disclosure of information submitted to the Commission pursuant to section 15(b) of the CPSA, 15 U.S.C. 2064(b).

(b) Criteria for disclosure. Under section 6(b)(5) the Commission shall not disclose to the public information which is identified as being submitted pursuant to section 15(b) or which is treated by the Commission staff as being submitted pursuant to section 15(b). Section 6(b)(5) also applies to information voluntarily submitted after a firm’s initial report to assist the Commission in its evaluation of the section 15 report. However, the Commission may disclose information submitted pursuant to section 15(b) in accordance with section 6(b)(1)–(3) if:

1. The Commission has issued a complaint under section 15 (c) or (d) of the CPSA alleging that such product presents a substantial product hazard; or
2. 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
3. The person who submitted the information under section 15(b) agrees to its public disclosure.

4. The Commission publishes a finding that the public health and safety requires public disclosure with a lesser period of notice than is required by section 6(b)(1).


§ 1101.62 Statutory exceptions to section 6(b)(5) requirements.

(a) Scope. The limitations established by section 6(b)(5) do not apply to the public disclosure of:

1. Information with respect to a consumer product which is the subject of an action brought under section 12 (see §1101.42);
2. Information with respect to a consumer product which the Commission has reasonable cause to believe is in violation of any consumer product safety rule or provision under the Consumer Product Safety Act (Pub. L. 92–573, 86 Stat. 1207, as amended (15 U.S.C. 2051, et seq.)) or similar rule or provision of any other act enforced by the Commission; or
3. Information in the course of or concerning a judicial proceeding (see §1101.45).


§ 1101.63 Information submitted pursuant to section 15(b) of the CPSA.

(a) Section 6(b)(5) applies only to information provided to the Commission by a manufacturer, distributor, or retailer which is identified by the manufacturer, distributor or retailer, or treated by the Commission staff as being submitted pursuant to section 15(b).

(b) Section 6(b)(5)’s limitation also applies to the portions of staff generated documents that contain, summarize or analyze such information submitted pursuant to section 15(b).