

§ 1102.24

(1) Identity of the submitter and/or the victim, including name, location, age, and gender;

(2) Consumer product, including serial or model number, date code, color, or size;

(3) Harm or risk of harm related to the use of the consumer product;

(4) Description of the incident related to use of the consumer product;

(5) Date or approximate date of the incident; and/or

(6) Category of submitter.

(c) *Timing.* To the extent practicable, the Commission will transmit a report of harm to the manufacturer or private labeler within five business days of submission of the completed report of harm. If the Commission cannot determine whom the manufacturer or private labeler is from the report of harm, or otherwise, then it will not post the report of harm on the Database but will maintain the report for internal agency use. Examples of circumstances that may arise that may make transmission of the report of harm impracticable within five business days include, but are not limited to:

(1) The manufacturer or private labeler is out of business with no identifiable successor;

(2) The submitter misidentified a manufacturer or private labeler;

(3) The report of harm contained inaccurate or insufficient contact information for a manufacturer or private labeler; or

(4) The Commission cannot locate valid contact information for a manufacturer or private labeler.

(d) *Method of transmission.* The Commission will use the method of transmission and contact information provided by the manufacturer or private labeler. The Commission will transmit reports of harm to a manufacturer or private labeler who has registered with the Commission as described in paragraph (f) of this section. If a manufacturer or private labeler has not registered with the Commission, the Commission will send reports of harm through the United States mail to the firm's principal place of business, unless the Commission selects another equally effective method of transmission.

16 CFR Ch. II (1-1-11 Edition)

(e) *Size limits of manufacturer comments.* The Commission may, in its discretion, limit the data size of comments, which may include attachments submitted, where such comments and attachments may negatively impact the technological or operational performance of the system.

(f) *Manufacturer registration.* Manufacturers and private labelers may register with the Commission to select a preferred method for receiving reports of harm that identify such firm as the manufacturer or private labeler. Manufacturers and private labelers that choose to register with the Commission must:

(1) Register with the Commission through a process identified for such registration;

(2) Provide and maintain updated contact information for the firm, including the name of the firm, title of a person to whom reports of harm should be directed, complete mailing address, telephone number, electronic mail address, and Web site address (if any); and

(3) Select a specified method to receive reports of harm that identify the firm as the manufacturer or private labeler of a consumer product.

(g) *Manufacturer comments.* A manufacturer or private labeler who receives a report of harm from the CPSC may comment on the information contained in such report of harm. The Commission, in its discretion, where it determines it is in the public interest, may choose not to publish a manufacturer comment in the Database. For example, it may not be in the public interest for the Commission to publish comments that, in the unlikely event, contain language reasonably described as lewd, lascivious, or obscene.

§ 1102.24 Designation of confidential information.

(a) For purposes of this section, "confidential information" is considered to be information that contains or relates to a trade secret or other matter referred to in 18 U.S.C. 1905 or that is subject to 5 U.S.C. 552(b)(4).

(b) A manufacturer or private labeler identified in a report of harm and who receives a report of harm from the CPSC may review such report of harm

Consumer Product Safety Commission

§ 1102.26

for confidential information and request that portions of the report of harm be designated as confidential information. Each requester seeking such a designation of confidential information bears the burden of proof and must:

(1) Specifically identify the exact portion(s) of the report of harm claimed to be confidential;

(2) State whether the information claimed to be confidential has ever been released in any manner to a person who was not an employee or in a confidential relationship with the company;

(3) State whether the information so specified is commonly known within the industry or is readily ascertainable by outside persons with a minimum of time and effort;

(4) If known, state the company's relationship with the victim and/or submitter of the report of harm and how the victim and/or submitter of the report of harm came to be in possession of such allegedly confidential information;

(5) State how the release of the information would be likely to cause substantial harm to the company's competitive position; and

(6) State whether the person submitting the request for treatment as confidential information is authorized to make claims of confidentiality on behalf of the person or organization concerned.

(c) *Manner of submission.* Requests for designation of confidential information may be submitted in the same manner as manufacturer comments as described in §1102.12(b). A request for designation of confidential treatment must be conspicuously marked.

(d) *Timing of submission.* In order to ensure that the allegedly confidential information is not placed in the database, a request for designation of confidential information must be received by the Commission in a timely manner prior to the 10th business day after the date on which the Commission transmits the report to the manufacturer or private labeler. If a request for confidential treatment is submitted in a timely fashion, the Commission will either make a determination on the claim prior to posting on the 10th busi-

ness day after transmittal to the manufacturer or, as a matter of policy, redact the allegedly confidential information from a report of harm before publication in the Database until it makes a determination regarding confidential treatment.

(e) *Assistance with defense.* No request to redact confidential information from a report of harm pursuant to 5 U.S.C. 552(b)(4) should be made by any person who does not intend in good faith, and so certifies in writing, to assist the Commission in the defense of any judicial proceeding that thereafter might be brought to compel the disclosure of information that the Commission has determined to be a trade secret or privileged or confidential commercial or financial information.

(f) *Commission determination of confidentiality.* If the Commission determines that information in a report of harm is confidential, the Commission shall:

(1) Notify the manufacturer or private labeler;

(2) Redact such confidential information in the report of harm; and

(3) Publish the report of harm in the Database without such confidential information.

(g) *Commission determination of no confidentiality.* If the Commission determines that a report of harm does not contain confidential information, the Commission shall:

(1) Notify the manufacturer or private labeler; and

(2) Publish the report of harm, if not already published, in the Database.

(h) *Removal of confidential information.* As stated at 6A(c)(1)(C)(iii) of the CPSA, to seek removal of alleged confidential information that has been published in the Database, a manufacturer or private labeler may bring an action in the district court of the United States in the district in which the complainant resides, or has its principal place of business, or in the U.S. District Court for the District of Columbia.

§ 1102.26 Determination of materially inaccurate information.

(a) For purposes of this section, the following definitions apply: