will publish manufacturer comments related to a report of harm transmitted to a manufacturer or private labeler in the Database if such manufacturer comment meets the following requirements:

(1) **Manufacturer comment relates to report of harm.** The manufacturer or private labeler’s comment must relate to information contained in a specific report of harm that identifies such manufacturer or private labeler and that is submitted for publication in the Database.

(2) **Unique identifier.** A manufacturer comment must state the unique identifier provided by the CPSC.

(3) **Verification.** A manufacturer or private labeler must verify that it has reviewed the report of harm and the comment related to the report of harm and that the information contained in the comment is true and accurate to the best of the firm’s knowledge, information, and belief.

(4) **Request for publication.** When a manufacturer or private labeler submits a comment regarding a report of harm, it may request that the Commission publish such comment in the Database. A manufacturer or private labeler must affirmatively request publication of the comment, and consent to such publication in the Database, for each comment submitted to the CPSC.

(d) **Information published.** Subject to §§1102.24 and 1102.26, the Commission will publish a manufacturer comment and the date of its submission to the CPSC in the Database if the comment meets the minimum requirements for publication as described in paragraph (c) of this section.

(e) **Information not published.** The Commission will not publish in the Database consents and verifications associated with a manufacturer comment.

§ 1102.14 Recall notices.

All information presented in a voluntary or mandatory recall notice that has been made available to the public shall be accessible and searchable in the Database.

§ 1102.16 Additional information.

In addition to reports of harm, manufacturer comments, and recall notices, the CPSC shall include in the Database any additional information it determines to be in the public interest, consistent with the requirements of section 6(a) and (b) of the CPSA.

Subpart C—Procedural Requirements

§ 1102.20 Transmission of reports of harm to the identified manufacturer or private labeler.

(a) **Information transmitted.** Except as provided in paragraphs (a)(1) through (a)(3) of this section, the Commission will transmit all information provided in a report of harm, provided such report meets the minimum requirements for publication in the Database, to the manufacturer or private labeler identified in a report of harm. The following information will not be transmitted to a manufacturer or private labeler:

(1) Name and contact information for the submitter of the report of harm, unless such submitter provides express written consent (for example, by checking a box on the report of harm) to provide such information to the manufacturer or private labeler;

(2) Photographs that could be used to identify a person; and

(3) Medical records, unless the person about whom such records pertain, or his or her parent, guardian, or appropriate legally authorized representative, consents to providing such records to the manufacturer or private labeler.

(b) **Limitation on use of contact information.** A manufacturer or private labeler who receives name and contact information for the submitter of a report of harm and/or a victim must not use or disseminate such information to any other party for any other purpose other than verification of information contained in a report of harm. Verification of information contained in a report of harm must not include activities such as sales, promotion, marketing, warranty, or any other commercial purpose. Verification of information contained in a report of harm may include verification of the: