

## Consumer Product Safety Commission

## § 1051.5

- 1051.5 Requirements and recommendations for petitions.
- 1051.6 Documents not considered petitions.
- 1051.7 Statement in support of or in opposition to petitions; Duty of petitioners to remain apprised of developments regarding petitions.
- 1051.8 Public hearings on petitions.
- 1051.9 Factors the Commission considers in granting or denying petitions.
- 1051.10 Granting petitions.
- 1051.11 Denial of petitions.

AUTHORITY: 5 U.S.C. 553(e), 5 U.S.C. 555(e).

SOURCE: 48 FR 57123, Dec. 28, 1983, unless otherwise noted.

### § 1051.1 Scope.

(a) This part establishes procedures for the submission and disposition of petitions for the issuance, amendment or revocation of rules under the Consumer Product Safety Act (CPSA) (15 U.S.C. 2051 *et seq.*) or other statutes administered by the Consumer Product Safety Commission.

(b) Persons filing petitions for rule-making shall follow as closely as possible the requirements and are encouraged to follow as closely as possible the recommendations for filing petitions under § 1051.5.

(c) Petitions regarding products regulated under the Federal Hazardous Substances Act (FHSA) (15 U.S.C. 1261 *et seq.*) are governed by existing Commission procedures at 16 CFR 1500.82. Petitions regarding the exemption of products regulated under the Poison Prevention Packaging Act of 1970 (PPPA) (15 U.S.C. 1471 *et seq.*) are governed by existing Commission procedures at 16 CFR part 1702. In addition, however, persons filing such petitions shall follow the requirements and are encouraged to follow the recommendations for filing petitions as set forth in § 1051.5.

[48 FR 57123, Dec. 28, 1983 as amended at 64 FR 48704, Sept. 8, 1999]

### § 1051.2 General.

(a) Any person may file with the Commission a petition requesting the Commission to begin a proceeding to issue, amend or revoke a regulation under any of the statutes it administers.

(b) A petition which addresses a risk of injury associated with a product which could be eliminated or reduced

to a sufficient extent by action taken under the Federal Hazardous Substances Act, the Poison Prevention Packaging Act of 1970, or the Flammable Fabrics Act may be considered by the Commission under those Acts. However, if the Commission finds by rule, in accordance with section 30(d) of the CPSA, as amended by Public Law 94-284, that it is in the public interest to regulate such risk of injury under the CPSA, it may do so. Upon determination by the Office of the General Counsel that a petition should be considered under one of these acts rather than the CPSA, the Office of the Secretary shall docket and process the petition under the appropriate act and inform the petitioner of this determination. Such docketing, however, shall not preclude the Commission from proceeding to regulate the product under the CPSA after making the necessary findings.

### § 1051.3 Place of filing.

A petition should be mailed to: Office of the Secretary, Consumer Product Safety Commission, Washington, DC 20207. Persons wishing to file a petition in person may do so in the Office of the Secretary, at 4330 East West Highway, Bethesda, Maryland.

[48 FR 57123, Dec. 28, 1983, as amended at 62 FR 46667, Sept. 4, 1997]

### § 1051.4 Time of filing.

For purposes of computing time periods under this part, a petition shall be considered filed when time-date stamped by the Office of the Secretary. A document is time-date stamped when it is received in the Office of the Secretary.

### § 1051.5 Requirements and recommendations for petitions.

(a) *Requirements.* To be considered a petition under this part, any request to issue, amend or revoke a rule shall meet the requirements of this paragraph (a). A petition shall:

- (1) Be written in the English language;
- (2) Contain the name and address of the petitioner;
- (3) Indicate the product (or products) regulated under the Consumer Product

## § 1051.6

Safety Act or other statute the Commission administers for which a rule is sought or for which there is an existing rule sought to be modified or revoked. (If the petition regards a procedural or other rule not involving a specific product, the type of rule involved must be indicated.)

(4) Set forth facts which establish the claim that the issuance, amendment, or revocation of the rule is necessary (for example, such facts may include personal experience; medical, engineering or injury data; or a research study); and

(5) Contain an explicit request to initiate Commission rulemaking and set forth a brief description of the substance of the proposed rule or amendment or revocation thereof which it is claimed should be issued by the Commission. (A general request for regulatory action which does not reasonably specify the type of action requested shall not be sufficient for purposes of this subsection.)

(b) *Recommendations.* The Commission encourages the submission of as much information as possible related to the petition. Thus, to assist the Commission in its evaluation of a petition, to the extent the information is known and available to the petitioner, the petitioner is encouraged to supply the following information or any other information relating to the petition. The petition will be considered by the Commission even if the petitioner is unable to supply the information recommended in this paragraph (b). However, as applicable, and to the extent possible, the petitioner is encouraged to:

(1) Describe the specific risk(s) of injury to which the petition is addressed, including the degree (severity) and the nature of the risk(s) of injury associated with the product and possible reasons for the existence of the risk of injury (for example, product defect, poor design, faulty workmanship, or intentional or unintentional misuse);

(2) State why a consumer product safety standard would not be feasible if the petition requests the issuance of a rule declaring the product to be a banned hazardous product; and

(3) Supply or reference any known documentation, engineering studies,

## 16 CFR Ch. II (1–1–07 Edition)

technical studies, reports of injuries, medical findings, legal analyses, economic analyses and environmental impact analyses relating to the petition.

(c) *Procedural recommendations.* The following are procedural recommendations to help the Commission in its consideration of petitions. The Commission requests, but does not require, that a petition filed under this part:

(1) Be typewritten,

(2) Include the word “petition” in a heading preceding the text,

(3) Specify what section of the statute administered by the Commission authorizes the requested rulemaking,

(4) Include the telephone number of the petitioner, and

(5) Be accompanied by at least five (5) copies of the petition.

### § 1051.6 Documents not considered petitions.

(a) A document filed with the Commission which addresses a topic or involves a product outside the jurisdiction of the Commission will not be considered to be a petition. After consultation with the Office of the General Counsel, the Office of the Secretary, if appropriate, will forward to the appropriate agency documents which address products or topics within the jurisdiction of other agencies. The Office of the Secretary shall notify the sender of the document that it has been forwarded to the appropriate agency.

(b) Any other documents filed with the Office of the Secretary that are determined by the Office of the General Counsel not to be petitions shall be evaluated for possible staff action. The Office of the General Counsel shall notify the writer of the manner in which the Commission staff is treating the document. If the writer has indicated an intention to petition the Commission, the Office of the General Counsel shall inform the writer of the procedure to be followed for petitioning.

### § 1051.7 Statement in support of or in opposition to petitions; Duty of petitioners to remain apprised of developments regarding petitions.

(a) Any person may file a statement with the Office of the Secretary in support of or in opposition to a petition