§ 1253.6 Certifying and nullifying an approval.

(a) An Enterprise shall certify, through an executive officer, as that term is defined by §1770.3(g) of this title, that any filing or supporting material submitted to FHFA pursuant to regulations in this part contains no material misrepresentations or omissions. FHFA may review and verify any information filed in connection with a Notice. If FHFA discovers a material misrepresentation or omission after the Director has rendered a decision on the filing, FHFA may nullify any approval or modify the terms, conditions, and limitations to such approval. For purposes of this paragraph, an Enterprise’s authority to offer a new product or engage in a new activity by reason of the Director’s not having made an explicit determination within the statutory time period constitutes an approval.

(b) Any person responsible for any material misrepresentation or omission in a submission or supporting materials may be subject to enforcement action and other penalties, including the following:

(1) Identification of the specific information for which confidential treatment is sought, and the specific Notice for which the information is being submitted;

(2) Explanation of the bases for the proposed confidential treatment including, but not limited to, why the information is "commercial or financial information obtained from a person and privileged or confidential" as that phrase is used in Exemption 4 of the Freedom of Information Act (FOIA), 5 U.S.C. 552(b)(4), and §1202.4(a)(4) of this chapter;

(3) Explanation of the relevance and necessity of the information to whether the Notice should be approved or denied;

(4) Explanation of how disclosure of the information would result in substantial harm to the competitive position of the Enterprise or commenter;

(5) Explanation of whether the information is available to the public and the extent of any previous disclosure to third parties;

(6) Justification of the time period during which the Enterprise or commenter asserts that the material should not be available for public disclosure; and

(7) Any other information that the Enterprise or commenter seeking confidential treatment believes may be useful in assessing whether its request for confidentiality should be granted.
§ 1253.7 Failure to comply.

(a) Unless the Director otherwise informs the Enterprise in writing, an Enterprise must cease offering a new product or engaging in a new activity immediately upon discovering or receiving notice from the Director that the Enterprise has—

(1) Offered a new product or commenced a new activity without submitting a Notice;

(2) Offered a new product or commenced a new activity after submitting a Notice but before approval is granted, and before the expiration of the time provided for the Director to make a determination under §§1253.3 and 1253.4;

(3) Offered a new product after the Director disapproved it; or

(4) Failed to adhere to any terms, conditions or limitations established by the Director in his or her approval of a new product or activity.

(b) Within five (5) business-days of the discovery or notice of any of the events described in paragraph (a) of this section, the Enterprise must provide the Director a written description of the failure or failures of controls that resulted in the offering of the new product or commencement of the new activity in contravention of this regulation, and the steps that the Enterprise has taken or will take to remediate the control failures. The Enterprise must provide the board of directors of the Enterprise and chief risk officer, internal audit, and compliance officer of the Enterprise with a copy of the written description on the same date the description is provided to the Director.

(c) In the event that the Enterprise elects to resubmit the Notice of a new product or new activity that was undertaken in contravention of this regulation, the resubmission must provide sufficient documentation of the effectiveness of the remediation efforts described in paragraph (b) of this section.

(d) Failure to comply with paragraphs (a) or (b) of this section above may result in FHFA’s taking enforcement action, including pursuant to 12 U.S.C. 4631 (orders to cease and desist), 12 U.S.C. 4632 (temporary orders to cease and desist), and 12 U.S.C. 4636 (civil money penalties).

§ 1253.8 Availability of new product to an Enterprise after it has been approved for the other Enterprise.

(a) If the Director approves a new product for one Enterprise or the new product is otherwise available to that Enterprise under §1253.4, the other Enterprise may also undertake that new product, subject to submitting a request to the Director in the form of a Notice under §1253.3 and approval by the Director.

(b) The Director may require such further information from the requesting Enterprise as he or she deems necessary to approve or deny the request. Approving the request does not require public notice and comment.

§ 1253.9 Preservation of authority.

(a) The Director’s exercise of his or her authority pursuant to the prior approval authority for products under section 1321 of the Safety and Soundness Act (12 U.S.C. 4541), and this regulation and other issuances in no way restricts—

(1) The safety and soundness authority of the Director over all new and existing products or activities; or

(2) The authority of the Director to review all new and existing products or activities to determine that such products or activities are consistent with the statutory mission of an Enterprise.

APPENDIX TO PART 1253—PRIOR APPROVAL FOR ENTERPRISE PRODUCTS—INSTRUCTIONS AND NOTICE OF NEW ACTIVITY FORM