Food and Drug Administration, HHS

(b) The Food and Drug Administration may refuse to consider any particular nonclinical laboratory study in support of an application for a research or marketing permit, if it finds that the study was not conducted in accordance with the good laboratory practice regulations set forth in this part, without disqualifying the testing facility that conducted the study or undertaking other regulatory action.

§ 58.217 Suspension or termination of a testing facility by a sponsor.

Termination of a testing facility by a sponsor is independent of, and neither in lieu of nor a precondition to, proceedings or actions authorized by this subpart. If a sponsor terminates or suspends a testing facility from further participation in a nonclinical laboratory study that is being conducted as part of any application for a research or marketing permit that has been submitted to any Center of the Food and Drug Administration (whether approved or not), it shall notify that Center in writing within 15 working days of the action; the notice shall include a statement of the reasons for such action. Suspension or termination of a testing facility by a sponsor does not relieve it of any obligation under any other applicable regulation to submit the results of the study to the Food and Drug Administration.


§ 58.219 Reinstatement of a disqualified testing facility.

A testing facility that has been disqualified may be reinstated as an acceptable source of nonclinical laboratory studies to be submitted to the Food and Drug Administration if the Commissioner determines, upon an evaluation of the submission of the testing facility, that the facility can adequately assure that it will conduct future nonclinical laboratory studies in compliance with the good laboratory practice regulations set forth in this part and, if any studies are currently being conducted, that the quality and integrity of such studies have not been seriously compromised. A disqualified testing facility that wishes to be so reinstated shall present in writing to the Commissioner reasons why it believes it should be reinstated and a detailed description of the corrective actions it has taken or intends to take to assure that the acts or omissions which led to its disqualification will not recur. The Commissioner may condition reinstatement upon the testing facility being found in compliance with the good laboratory practice regulations upon an inspection. If a testing facility is reinstated, the Commissioner shall so notify the testing facility and all organizations and persons who were notified, under §58.213 of the disqualification of the testing facility. A determination that a testing facility has been reinstated is disclosable to the public under part 20 of this chapter.

PART 60—PATENT TERM RESTORATION

Subpart A—General Provisions

Sec.
60.1 Scope.
60.2 Purpose.
60.3 Definitions.

Subpart B—Eligibility Assistance

60.10 FDA assistance on eligibility.

Subpart C—Regulatory Review Period Determinations

60.20 FDA action on regulatory review period determinations.
60.22 Regulatory review period determinations.
60.24 Revision of regulatory review period determinations.
60.26 Final action on regulatory review period determinations.
60.28 Time frame for determining regulatory review periods.

Subpart D—Due Diligence Petitions

60.30 Filing, format, and content of petitions.
60.32 Applicant response to petition.
60.34 FDA action on petitions.
60.36 Standard of due diligence.

Subpart E—Due Diligence Hearings

60.40 Request for hearing.
60.42 Notice of hearing.
60.44 Hearing procedures.
60.46 Administrative decision.


319