

(d) Review

Not later than 180 days after receipt of an ICCVAM test recommendation, a Federal agency carrying out a program described in subsection (a) of this section shall review such recommendation and notify the ICCVAM in writing of its findings.

(e) Recommendation adoption

Each Federal agency carrying out a program described in subsection (a) of this section, or its specific regulatory unit or units, shall adopt the ICCVAM test recommendation unless such Federal agency determines that—

(1) the ICCVAM test recommendation is not adequate in terms of biological relevance for the regulatory goal authorized by that agency, or mandated by Congress;

(2) the ICCVAM test recommendation does not generate data, in an amount and of a scientific value that is at least equivalent to the data generated prior to such recommendation, for the appropriate hazard identification, dose-response assessment, or risk assessment purposes as the current test method recommended or required by that agency;

(3) the agency does not employ, recommend, or require testing for that class of chemical or for the recommended test endpoint; or

(4) the ICCVAM test recommendation is unacceptable for satisfactorily fulfilling the test needs for that particular agency and its respective congressional mandate.

(Pub. L. 106-545, § 4, Dec. 19, 2000, 114 Stat. 2723.)

CODIFICATION

Section was enacted as part of the ICCVAM Authorization Act of 2000, and not as part of the Public Health Service Act which comprises this chapter.

§ 2851-5. Application**(a) Application**

Sections 2851-2 to 2851-5 of this title shall not apply to research, including research performed using biotechnology techniques, or research related to the causes, diagnosis, treatment, control, or prevention of physical or mental diseases or impairments of humans or animals.

(b) Use of test methods

Nothing in sections 2851-2 to 2851-5 of this title shall prevent a Federal agency from retaining final authority for incorporating the test methods recommended by the ICCVAM in the manner determined to be appropriate by such Federal agency or regulatory body.

(c) Limitation

Nothing in sections 2851-2 to 2851-5 of this title shall be construed to require a manufacturer that is currently not required to perform animal testing to perform such tests. Nothing in sections 2851-2 to 2851-5 of this title shall be construed to require a manufacturer to perform redundant endpoint specific testing.

(d) Submission of tests and data

Nothing in sections 2851-2 to 2851-5 of this title precludes a party from submitting a test method or scientific data directly to a Federal agency for use in a regulatory program.

(Pub. L. 106-545, § 5, Dec. 19, 2000, 114 Stat. 2724.)

CODIFICATION

Section was enacted as part of the ICCVAM Authorization Act of 2000, and not as part of the Public Health Service Act which comprises this chapter.

§ 2851-6. Methods of controlling certain insect and vermin populations

The Director of the Institute shall conduct or support research to identify or develop methods of controlling insect and vermin populations that transmit to humans diseases that have significant adverse health consequences.

(July 1, 1944, ch. 373, title IV, § 463B, as added Pub. L. 108-75, § 3, Aug. 15, 2003, 117 Stat. 902.)

SUBPART 13—NATIONAL INSTITUTE ON DEAFNESS
AND OTHER COMMUNICATION DISORDERS**§ 285m. Purpose of Institute**

The general purpose of the National Institute on Deafness and Other Communication Disorders (hereafter referred to in this subpart as the “Institute”) is the conduct and support of research and training, the dissemination of health information, and other programs with respect to disorders of hearing and other communication processes, including diseases affecting hearing, balance, voice, speech, language, taste, and smell.

(July 1, 1944, ch. 373, title IV, § 464, as added Pub. L. 100-553, § 2(4), Oct. 28, 1988, 102 Stat. 2769, and Pub. L. 100-607, title I, § 101(4), Nov. 4, 1988, 102 Stat. 3049; amended Pub. L. 100-690, title II, § 2613(b)(2), Nov. 18, 1988, 102 Stat. 4238.)

CODIFICATION

Pub. L. 100-553 and Pub. L. 100-607 contained identical provisions enacting this section. See 1988 Amendment note below.

AMENDMENTS

1988—Pub. L. 100-690 amended this section to read as if the amendments made by Pub. L. 100-607, which enacted this section, had not been enacted. See Codification note above.

SHORT TITLE OF 1988 AMENDMENT

For short title of Pub. L. 100-553 which enacted this subpart and amended sections 281 and 285j of this title as the “National Deafness and Other Communication Disorders Act of 1988”, see section 1 of Pub. L. 100-553, set out as a note under section 201 of this title.

EFFECT OF ENACTMENT OF SIMILAR PROVISIONS

Section 2613(b) of Pub. L. 100-690 provided that:

“(1) Paragraphs (2) and (3) shall take effect immediately after the enactment of both the bill, S. 1727, of the One Hundredth Congress [Pub. L. 100-553, approved Oct. 28, 1988], and the Health Omnibus Programs Extension of 1988 [Pub. L. 100-607, approved Nov. 4, 1988].

“(2)(A) The provisions of the Public Health Service Act referred to in subparagraph (B), as similarly amended by the enactment of the bill, S. 1727, of the One Hundredth Congress, by subtitle A of title I of the Health Omnibus Programs Extension of 1988, and by subsection (a)(1) of this section, are amended to read as if the amendments made by such subtitle A and such subsection (a)(1) had not been enacted.

“(B) The provisions of the Public Health Service Act referred to in subparagraph (A) are—

“(A) sections 401(b)(1) and 457 [sections 281(b)(1) and 285j of this title];