

119TH CONGRESS
2D SESSION

S. 4472

AN ACT

To amend the Accelerating Access to Critical Therapies for ALS Act to reauthorize the provisions of such Act through fiscal year 2031, and for other purposes.

1 *Be it enacted by the Senate and House of Representa-*
2 *tives of the United States of America in Congress assembled,*

1 **SECTION 1. SHORT TITLE.**

2 This Act may be cited as the “Accelerating Access
3 to Critical Therapies for ALS Reauthorization Act of
4 2026”.

5 **SEC. 2. REAUTHORIZATION OF ACCELERATING ACCESS TO**
6 **CRITICAL THERAPIES FOR ALS ACT.**

7 (a) IN GENERAL.—Section 7 of the Accelerating Ac-
8 cess to Critical Therapies for ALS Act (Public Law 117–
9 79) is amended by striking “2022 through 2026” and in-
10 serting “2027 through 2031”.

11 (b) GRANTS FOR ALS RESEARCH.—Section 2(f) of
12 the Accelerating Access to Critical Therapies for ALS Act
13 (21 U.S.C. 360ee note) is amended by striking “2026”
14 and inserting “2031”.

15 **SEC. 3. IMPROVEMENTS TO PROGRAM FOR GRANTS FOR**
16 **RESEARCH ON THERAPIES FOR ALS.**

17 Section 2 of the Accelerating Access to Critical
18 Therapies for ALS Act (21 U.S.C. 360ee note) is amend-
19 ed—

20 (1) in subsection (a), by inserting “(referred to
21 in this section as ‘expanded access grants’)” before
22 the period at the end of the first sentence;

23 (2) in subsection (b)—

24 (A) by striking “(b) APPLICATION—” and
25 all that follows through “A participating” in
26 paragraph (1) and inserting the following:

1 “(b) APPLICATION.—A participating”;

2 (B) by redesignating paragraphs (2) and
3 (3) as paragraphs (1) and (2) respectively;

4 (C) in the matter preceding paragraph (1),
5 as so redesignated, by striking the period at the
6 end and inserting “including—”;

7 (D) by amending paragraph (1), as so re-
8 designated, to read as follows:

9 “(1) a description of how data generated
10 through the proposed expanded access grant will be
11 used to support research or development related to
12 the prevention, diagnosis, mitigation, treatment, or
13 cure of amyotrophic lateral sclerosis;”;

14 (E) in paragraph (2), as so redesignated—

15 (i) by striking “NONINTERFERENCE
16 WITH CLINICAL TRIALS—” and all that fol-
17 lows through “shall include”;

18 (ii) by striking “program” and insert-
19 ing “grant”; and

20 (iii) by striking the period at the end
21 and inserting “; and”; and

22 (F) by adding at the end the following:

23 “(3) an assurance that such entity will prompt-
24 ly report to the Secretary available safety data from
25 any ongoing clinical trial of the investigational drug

1 as set forth in the terms and conditions of the
 2 grant.”;

3 (3) in subsection (c)—

4 (A) by redesignating subparagraphs (A)
 5 and (B) of paragraph (2) as clauses (i) and (ii),
 6 respectively, and adjusting the margins accord-
 7 ingly;

8 (B) by redesignating paragraphs (1)
 9 through (3) as subparagraphs (A) through (C),
 10 respectively, and adjusting the margins accord-
 11 ingly;

12 (C) in subparagraph (C), as so redesign-
 13 nated, by striking the period at the end and in-
 14 serting “; and”;

15 (D) in the matter preceding subparagraph
 16 (A), as so redesignated, by striking “this sec-
 17 tion, confirm that—” and inserting the fol-
 18 lowing: “this section—

19 “(1) confirm that—”; and

20 (E) by adding at the end the following:

21 “(2) in the case of a renewal of such a grant,
 22 request from the sponsor of the investigational new
 23 drug application involved, and assess, the enroll-
 24 ment, safety, and any available efficacy data of the

1 drug related to the prevention, diagnosis, mitigation,
2 treatment, or cure of amyotrophic lateral sclerosis.”;

3 (4) in subsection (d)(1), by striking “request
4 described in subsection (a)” and inserting “grant”;
5 and

6 (5) in subsection (e)—

7 (A) in paragraph (2), by inserting “, and
8 that begins enrollment within a timeframe as
9 determined by the Secretary through the terms
10 and conditions of the grant” before the period
11 at the end; and

12 (B) by adding at the end the following:

13 “(4) The term ‘phase 3 clinical trial’ includes a
14 phase 2/3 combined trial and a planned phase 3 clin-
15 ical trial that is not yet enrolling participants.”.

16 **SEC. 4. REPORT ON ALS AND OTHER RARE**
17 **NEURODEGENERATIVE DISEASE ACTION**
18 **PLANS.**

19 Section 4 of the Accelerating Access to Critical
20 Therapies for ALS Act (21 U.S.C. 360aa note) is amend-
21 ed—

22 (1) in the section heading, by striking “**ALS**
23 **AND OTHER**” and inserting “**FDA**”;

24 (2) in subsection (a), in the matter preceding
25 paragraph (1)—

1 (A) by inserting “and not later than 1 year
 2 after the date of enactment of the Accelerating
 3 Access to Critical Therapies for ALS Reauthor-
 4 ization Act of 2026 and every 5 years there-
 5 after,” after “this Act,”; and

6 (B) by inserting “develop, or update, as
 7 applicable, and” before “publish on”;

8 (3) in subsection (b)—

9 (A) in the matter preceding paragraph (1),
 10 by striking “initial”;

11 (B) in paragraph (2)—

12 (i) in subparagraph (A), by inserting
 13 “of relevant investigational new drug appli-
 14 cations” after “sponsors”;

15 (ii) in subparagraph (C) by inserting
 16 “for the prevention, diagnosis, mitigation,
 17 treatment, or cure of rare
 18 neurodegenerative diseases” before the
 19 semicolon; and

20 (iii) in subparagraph (D), by striking
 21 “; and” and inserting a semicolon;

22 (C) in paragraph (3), by striking the pe-
 23 riod at the end and inserting “; and”; and

24 (D) by adding at the end the following:

1 “(4) for each action plan published after the
 2 date of enactment of the Accelerating Access to Crit-
 3 ical Therapies for ALS Reauthorization Act of 2026,
 4 include a description of—

5 “(A) previous actions taken by the Food
 6 and Drug Administration to implement the pre-
 7 vious action plan published under subsection
 8 (a);

9 “(B) any other planned actions to imple-
 10 ment such action plan; and

11 “(C) any barriers to implementing such ac-
 12 tion plan and related recommendations, which
 13 may include an estimate of resources necessary
 14 to address such barriers.”.

15 **SEC. 5. REPORTS.**

16 Section 6 of the Accelerating Access to Critical
 17 Therapies for ALS Act (Public Law 117–79) is amend-
 18 ed—

19 (1) in the heading, by striking “**GAO REPORT**”
 20 and inserting “**REPORTS**”;

21 (2) by striking “Not later than” and inserting
 22 the following:

23 “(a) GAO REPORT.—Not later than”;

24 (3) in the matter preceding paragraph (1) of
 25 subsection (a), as so designated, by striking “this

1 Act” and inserting “the Accelerating Access to Crit-
2 ical Therapies for ALS Reauthorization Act of
3 2026”; and

4 (4) by adding at the end the following:

5 “(b) HHS REPORT.—Not later than 4 years after the
6 date of enactment of the Accelerating Access to Critical
7 Therapies for ALS Reauthorization Act of 2026, the Sec-
8 retary of Health and Human Services shall, in a manner
9 that does not duplicate the information described in the
10 action plan published pursuant to section 4, submit to the
11 Committee on Health, Education, Labor, and Pensions of
12 the Senate and the Committee on Energy and Commerce
13 of the House of Representatives a report assessing the ef-
14 fectiveness of the activities carried out under sections 2,
15 3, and 5 and making recommendations to improve such
16 activities.”.

17 **SEC. 6. TECHNICAL AMENDMENTS.**

18 Section 3 of the Accelerating Access to Critical
19 Therapies for ALS Act (42 U.S.C. 280g–7b) is amend-
20 ed—

21 (1) in subsection (a), in the matter preceding
22 paragraph (1), by striking “amytrophic” and insert-
23 ing “amyotrophic”; and

- 1 (2) in subsection (b)(3)(A)(iii), by striking “ra-
2 tional”.

Passed the Senate August 4, 2026.

Attest:

Secretary.

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