117TH CONGRESS 1ST SESSION H. R. 482

AN ACT

To amend the Public Health Service Act to reauthorize certain programs under part A of title XI of such Act relating to genetic diseases, 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 "Newborn Screening3 Saves Lives Reauthorization Act of 2021".

4 SEC. 2. IMPROVED NEWBORN AND CHILD SCREENING AND 5 FOLLOW-UP FOR HERITABLE DISORDERS.

6 (a) PURPOSES.—Section 1109(a) of the Public
7 Health Service Act (42 U.S.C. 300b-8(a)) is amended—
8 (1) in paragraph (1), by striking "enhance, im9 prove or" and inserting "facilitate, enhance, im10 prove, or";

(2) by amending paragraph (3) to read as fol-lows:

13 "(3) to develop, and deliver to parents, families,
14 and patient advocacy and support groups, edu15 cational programs that—

"(A) address newborn screening counseling, testing (including newborn screening
pilot studies), follow-up, treatment, specialty
services, and long-term care;

20 "(B) assess the target audience's current
21 knowledge, incorporate health communications
22 strategies, and measure impact; and

23 "(C) are at appropriate literacy levels;";24 and

(3) in paragraph (4)—

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1	(A) by striking "followup" and inserting
2	"follow-up"; and
3	(B) by inserting before the semicolon at
4	the end the following: ", including re-engaging
5	patients who have not received recommended
6	follow-up services and supports".
7	(b) Approval Factors.—Section 1109(c) of the
8	Public Health Service Act (42 U.S.C. 300b-8(c)) is
9	amended—
10	(1) by striking "or will use" and inserting "will
11	use"; and
12	(2) by inserting ", or will use amounts received
13	under such grant to enhance capacity and infra-
14	structure to facilitate the adoption of," before "the
15	guidelines and recommendations".
16	SEC. 3. ADVISORY COMMITTEE ON HERITABLE DISORDERS
17	IN NEWBORNS AND CHILDREN.
18	Section 1111 of the Public Health Service Act (42)
19	U.S.C. 300b–10) is amended—
20	(1) in subsection (b)—
21	(A) in paragraph (5), by inserting "and
22	adopt process improvements'' after "take ap-
23	propriate steps";
24	(B) in paragraph (7) by striking "and" at
25	the end;

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1	(C) by redesignating paragraph (8) as
2	paragraph (9);
3	(D) by inserting after paragraph (7) the
4	following:
5	"(8) develop, maintain, and publish on a pub-
6	licly accessible website consumer-friendly materials
7	detailing—
8	"(A) the uniform screening panel nomina-
9	tion process, including data requirements,
10	standards, and the use of international data in
11	nomination submissions; and
12	"(B) the process for obtaining technical as-
13	sistance for submitting nominations to the uni-
14	form screening panel and detailing the in-
15	stances in which the provision of technical as-
16	sistance would introduce a conflict of interest
17	for members of the Advisory Committee; and";
18	(E) in paragraph (9), as redesignated—
19	(i) by redesignating subparagraphs
20	(K) and (L) as subparagraphs (L) and
21	(M), respectively; and
22	(ii) by inserting after subparagraph
23	(J) the following:
24	"(K) the appropriate and recommended
25	use of safe and effective genetic testing by

health care professionals in newborns and chil-	
dren with an initial diagnosis of a disease or	
condition characterized by a variety of genetic	
causes and manifestations;"; and	
(2) in subsection (g)—	
(A) in paragraph (1) by striking "2019"	
and inserting "2026"; and	
(B) in paragraph (2) by striking "2019"	
and inserting "2026".	
SEC. 4. CLEARINGHOUSE OF NEWBORN SCREENING INFOR-	
MATION.	
Section 1112(c) of the Public Health Service Act (42	
U.S.C. 300b–11(c)) is amended by striking "and supple-	
ment, not supplant, existing information sharing efforts"	
and inserting "and complement other Federal newborn	

SEC. 5. LABORATORY QUALITY AND SURVEILLANCE.

screening information sharing activities".

Section 1113 of the Public Health Service Act (42) U.S.C. 300b–12) is amended—

(1) in subsection (a)—

(A) in paragraph (1)—

(i) by striking "performance evaluation services," and inserting "development of new screening tests,"; and

(ii) by striking "and" at the end;

1	(B) in paragraph (2)—
2	(i) by striking "performance test ma-
3	terials" and inserting "test performance
4	materials"; and
5	(ii) by striking the period at the end
6	and inserting "; and"; and
7	(C) by adding at the end the following:
8	"(3) performance evaluation services to enhance
9	disease detection, including the development of tools,
10	resources, and infrastructure to improve data anal-
11	ysis, test result interpretation, data harmonization,
12	and dissemination of laboratory best practices."; and
13	(2) in subsection (b) to read as follows:
14	"(b) SURVEILLANCE ACTIVITIES.—The Secretary,
15	acting through the Director of the Centers for Disease
16	Control and Prevention, and taking into consideration the
17	expertise of the Advisory Committee on Heritable Dis-
18	orders in Newborns and Children established under sec-
19	tion 1111, shall provide for the coordination of national
20	surveillance activities, including—
21	((1) standardizing data collection and reporting
22	through the use of electronic and other forms of
23	health records to achieve real-time data for tracking
24	and monitoring the newborn screening system, from

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1	the initial positive screen through diagnosis and
2	long-term care management; and
3	"(2) by promoting data sharing linkages be-
4	tween State newborn screening programs and State-
5	based birth defects and developmental disabilities
6	surveillance programs to help families connect with
7	services to assist in evaluating long-term outcomes.".
8	SEC. 6. HUNTER KELLY RESEARCH PROGRAM.
9	Section 1116 of the Public Health Service Act (42)
10	U.S.C. 300b–15) is amended—
11	(1) in subsection $(a)(1)$ —
12	(A) by striking "may" and inserting
13	"shall"; and
14	(B) in subparagraph (D)—
15	(i) by inserting ", or with a high prob-
16	ability of being recommended by," after
16 17	
	ability of being recommended by," after
17	ability of being recommended by," after "recommended by"; and
17 18	ability of being recommended by," after "recommended by"; and (ii) by striking "that screenings are
17 18 19	ability of being recommended by," after "recommended by"; and (ii) by striking "that screenings are ready for nationwide implementation" and
17 18 19 20	ability of being recommended by," after "recommended by"; and (ii) by striking "that screenings are ready for nationwide implementation" and inserting "that reliable newborn screening
 17 18 19 20 21 	ability of being recommended by," after "recommended by"; and (ii) by striking "that screenings are ready for nationwide implementation" and inserting "that reliable newborn screening technologies are piloted and ready for
 17 18 19 20 21 22 	ability of being recommended by," after "recommended by"; and (ii) by striking "that screenings are ready for nationwide implementation" and inserting "that reliable newborn screening technologies are piloted and ready for use"; and

1	shall ensure that entities receiving funding through the
2	program will provide assurances, as practicable, that such
3	entities will work in consultation with State departments
4	of health, as appropriate.".
5	SEC. 7. AUTHORIZATION OF APPROPRIATIONS FOR NEW-
6	BORN SCREENING PROGRAMS AND ACTIVI-
7	TIES.
8	Section 1117 of the Public Health Service Act (42)
9	U.S.C. 300b–16) is amended—
10	(1) in paragraph (1) —
11	(A) by striking "\$11,900,000" and insert-
12	ing ''\$31,000,000'';
13	(B) by striking "2015" and inserting
14	"2022"; and
15	(C) by striking "2019" and inserting
16	"2026"; and
17	(2) in paragraph (2) —
18	(A) by striking "\$8,000,000" and inserting
19	``\$29,650,000'';
20	(B) by striking "2015" and inserting
21	"2022"; and
22	(C) by striking "2019" and inserting
23	<i>"2026"</i> .

1 SEC. 8. INSTITUTIONAL REVIEW BOARDS; ETHICS GUID 2 ANCE PROGRAM.

3 Section 12 of the Newborn Screening Saves Lives Re4 authorization Act of 2014 (42 U.S.C. 289 note) is amend5 ed to read as follows:

6 "SEC. 12. INSTITUTIONAL REVIEW BOARDS; ETHICS GUID7 ANCE PROGRAM.

8 "Research on nonidentified newborn dried blood spots 9 shall be considered secondary research (as that term is 10 defined in section 46.104(d)(4) of title 45, Code of Federal 11 Regulations (or successor regulations)) with nonidentified 12 biospecimens for purposes of federally funded research 13 conducted pursuant to the Public Health Service Act (42 14 U.S.C. 200 et seq.).".

15 SEC. 9. NAM REPORT ON THE MODERNIZATION OF NEW16 BORN SCREENING.

17 (a) STUDY.—Not later than 60 days after the date 18 of the enactment of this Act, the Secretary of Health and 19 Human Services shall seek to enter into an agreement with the National Academy of Medicine (in this section 20 21 referred to as "NAM") (or if NAM declines to enter into such an agreement, another appropriate entity) under 22 23 which NAM, or such other appropriate entity, agrees to 24 conduct a study on the following:

(1) The uniform screening panel review and
recommendation processes to identify factors that
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impact decisions to add new conditions to the uniform screening panel, to describe challenges posed
by newly nominated conditions, including low-incidence diseases, late onset variants, and new treatments without long-term efficacy data.

6 (2) The barriers that preclude States from add-7 ing new uniform screening panel conditions to their 8 State screening panels with recommendations on re-9 sources needed to help States implement uniform 10 screening panel recommendations.

(3) The current state of federally and privately
funded newborn screening research with recommendations for optimizing the capacity of this research, including piloting multiple prospective conditions at once and addressing rare disease questions.

16 (4) New and emerging technologies that would
17 permit screening for new categories of disorders, or
18 would make current screening more effective, more
19 efficient, or less expensive.

20 (5) Technological and other infrastructure
21 needs to improve timeliness of diagnosis and short22 and long-term follow-up for infants identified
23 through newborn screening and improve public
24 health surveillance.

(6) Current and future communication and edu cational needs for priority stakeholders and the pub lic to promote understanding and knowledge of a
 modernized newborn screening system with an em phasis on evolving communication channels and mes saging.

7 (7) The extent to which newborn screening 8 yields better data on the disease prevalence for 9 screened conditions and improves long-term out-10 comes for those identified through newborn screen-11 ing, including existing systems supporting such data 12 collection and recommendations for systems that 13 would allow for improved data collection.

14 (8) The impact on newborn morbidity and mor15 tality in States that adopt newborn screening tests
16 included on the uniform panel.

(b) PUBLIC STAKEHOLDER MEETING.—In the course
of completing the study described in subsection (a), NAM
or such other appropriate entity shall hold not less than
one public meeting to obtain stakeholder input on the topics of such study.

(c) REPORT.—Not later than 18 months after the effective date of the agreement under subsection (a), such
agreement shall require NAM, or such other appropriate
entity, to submit to the Secretary of Health and Human

1	Services and the appropriate committees of jurisdiction of
2	Congress a report containing—
3	(1) the results of the study conducted under
4	subsection (a);
5	(2) recommendations to modernize the proc-
6	esses described in subsection $(a)(1)$; and
7	(3) recommendations for such legislative and
8	administrative action as NAM, or such other appro-
9	priate entity, determines appropriate.
10	(d) Authorization of Appropriations.—There is
11	authorized to be appropriated \$2,000,000 for the period
12	of fiscal years 2022 and 2023 to carry out this section.
	Passed the House of Representatives June 23, 2021.
	Attest:

Clerk.

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AN ACT

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