

117TH CONGRESS
1ST SESSION

H. R. 5989

To require the Secretary of Health and Human Services to conduct a demonstration program to genetic and genomic testing for certain children, to provide for a study by the National Academy of Medicine on the use of such testing to improve health care, and for other purposes.

IN THE HOUSE OF REPRESENTATIVES

NOVEMBER 16, 2021

Mr. SWALWELL (for himself, Mr. PETERS, and Mr. EMMER) introduced the following bill; which was referred to the Committee on Energy and Commerce, and in addition to the Committee on Ways and Means, for a period to be subsequently determined by the Speaker, in each case for consideration of such provisions as fall within the jurisdiction of the committee concerned

A BILL

To require the Secretary of Health and Human Services to conduct a demonstration program to genetic and genomic testing for certain children, to provide for a study by the National Academy of Medicine on the use of such testing to improve health care, 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 “Precision Medicine An-
3 swers for Kids Today Act”.

4 **SEC. 2. CENTERS FOR MEDICARE & MEDICAID SERVICES**
5 **GUIDANCE ON THE EARLY AND PERIODIC**
6 **SCREENING, DIAGNOSTIC, AND TREATMENT**
7 **BENEFIT.**

8 Not later than 6 months after the date of enactment
9 of this Act, the Centers for Medicare & Medicaid Services
10 shall issue guidance to States on authority and require-
11 ments under the Medicaid program under title XIX of the
12 Social Security Act to provide medically necessary health
13 care that falls within the scope of services specified under
14 section 1905(r) of the Social Security Act (42 U.S.C.
15 1396d(r)) to a child, regardless of whether the service is
16 available for adults under the State plan (or waiver of
17 such plan) under such title. The guidance shall—

18 (1) include technical and educational assistance
19 on how to increase the frequency of coverage under
20 the State plan (or waiver) pursuant to paragraphs
21 (4) and (16) of section 1905(a) of such Act (42
22 U.S.C. 1396d(a)) for genetic and genomic testing di-
23 agnostic services, including whole exome sequencing,
24 whole genome sequencing, and gene panels when rec-
25 ommended by a qualified treating provider as a first-

1 or second-tier test for pediatric patients, including
2 those who—

3 (A) have a positive result from a newborn
4 screening program;

5 (B) have one or more neurodevelopmental
6 or congenital anomalies;

7 (C) are experiencing developmental delay
8 or intellectual disability;

9 (D) are having seizures;

10 (E) have been referred or admitted to a
11 pediatric or neonatal intensive care unit for a
12 chronic or undiagnosed disease;

13 (F) have been seen by at least one medical
14 specialist for such chronic or undiagnosed dis-
15 ease; or

16 (G) are suspected by at least one
17 healthcare provider to have a neonatal- or pedi-
18 atric-onset genetic disease;

19 (2) provide education and support to providers
20 to minimize denials of claims for medical assistance
21 under the State plan under title XIX of the Social
22 Security Act resulting from deficient or inadequate
23 paperwork; and

24 (3) ensure that providers and Medicaid-eligible
25 children and the families are aware of the Early and

1 Periodic Screening, Diagnostic and Treatment Ben-
2 efit under title XIX of the Social Security Act and
3 have access to required screenings and necessary
4 treatment services.

5 **SEC. 3. DEMONSTRATION PROGRAM TO PROVIDE GENETIC**
6 **AND GENOMIC TESTING FOR CERTAIN CHIL-**
7 **DREN.**

8 (a) IN GENERAL.—The Secretary of Health and
9 Human Services shall enter into agreements with up to
10 15 States submitting applications under subsection (c) for
11 the purpose of conducting, in accordance with this section,
12 3-year demonstration projects under section 1115 of the
13 Social Security Act (42 U.S.C. 1315) in such States dur-
14 ing the 3-year period beginning on the first date of the
15 first fiscal quarter than begins on or after the date of the
16 enactment of this section to test and evaluate the provision
17 of medical assistance under the State plans under title
18 XIX of such Act (or waivers of such plans) to eligible indi-
19 viduals for purposes of providing such individuals with ge-
20 netic and genomic testing.

21 (b) DEMONSTRATION PROJECT PAYMENT REQUIRE-
22 MENTS.—Under each demonstration project under this
23 section conducted by a State, the following shall apply:

24 (1) The State shall provide a health care pro-
25 vider (as defined by the State) with payments for

1 the provision of genetic and genomic testing to any
2 eligible individual. Payments made to a health care
3 provider for such services shall be treated as medical
4 assistance for purposes of section 1903(a) of the So-
5 cial Security Act (42 U.S.C. 1396b(a)), except that
6 the Federal medical assistance percentage applicable
7 to such payments shall be equal to 100 percent.

8 (2) The State shall specify the methodology the
9 State will use for determining payment for the provi-
10 sion of genetic and genomic testing. Such method-
11 ology for determining payment shall be established
12 consistent with section 1902(a)(30)(A) of such Act
13 (42 U.S.C. 1396a(a)(30)(A)).

14 (c) APPLICATIONS.—

15 (1) IN GENERAL.—A State desiring to enter
16 into an agreement under subsection (a) with the
17 Secretary for conducting a demonstration project
18 shall submit to the Secretary an application, in ac-
19 cordance with such form and manner, and applica-
20 tion priorities, as specified by the Secretary and that
21 at a minimum includes the following:

22 (A) An explanation of how and the extent
23 to which genetic and genomic testing under the
24 demonstration project of the State will provide

1 information and data on how such services im-
2 prove the diagnosis of eligible individuals.

3 (B) An explanation of how and the extent
4 to which coverage under the State plan (or
5 waiver) pursuant to the demonstration project
6 will increase the use of genetic and genomic
7 testing that may increase the use of genetic and
8 genomic testing that may improve clinical out-
9 comes for eligible individuals.

10 (C) Procedures for referring any eligible
11 individual who seeks or needs treatment in a
12 hospital emergency department to a health care
13 provider who is qualified (as determined by the
14 State) to provide genetic and genomic testing.

15 (D) An explanation of how genetic and
16 genomic testing may improve health outcomes
17 for all populations in the State, including—

18 (i) individuals with a rare genetic dis-
19 ease, including a metabolic disease,
20 neurologic disorders, or hereditary cancer
21 testing in the presence of a suspected or
22 confirmed cancer diagnosis; and

23 (ii) special populations, including in-
24 fants and children who are critically ill
25 (non-infectious and non-trauma) patients,

1 transplant patients, individuals with car-
2 diac disease, and individuals with, or who
3 have a family history of, a birth defect or
4 developmental disability.

5 (2) PREFERENCES IN CONSIDERING APPLICA-
6 TIONS.—In considering applications submitted under
7 paragraph (1), the Secretary of Health and Human
8 Services shall give preference to States that can
9 demonstrate underutilization of genetic and genomic
10 sequencing clinical services (with priority given to
11 States that do not cover whole-genome sequencing or
12 do not cover the majority of genetic and genomic
13 clinical services) in pediatric populations under the
14 State plan under title XIX of the Social Security Act
15 (or waiver of such plan).

16 (d) TECHNICAL ASSISTANCE.—The Secretary of
17 Health and Human Services shall provide technical assist-
18 ance to assist States in planning and designing the dem-
19 onstration project for purposes of applying for conducting
20 such project under this section.

21 (e) REPORTS BY STATES.—Not later than one year
22 after the date on which a State enters into an agreement
23 under subsection (a) with the Secretary for conducting a
24 demonstration project, the State shall submit a report to
25 the Administrator of the Centers for Medicare & Medicaid

1 Services and the Administrator of the Health Resources
2 and Services Administration on the extent to which genetic
3 and genomic testing improved outcomes and reduced
4 health disparities. Such report shall include information
5 on the number of patients receiving genetic and genomic
6 testing, the types of services provided, and such other in-
7 formation as the Secretary shall prescribe.

8 (f) REPORTS BY HEALTH CARE PROVIDERS.—As a
9 condition for receiving payment for genetic and genomic
10 testing provided to an eligible individual under a dem-
11 onstration project conducted by a State under this section,
12 a health care provider shall report to the State, in accord-
13 ance with such requirements as the Secretary shall specify,
14 on all applicable measures for determining the quality and
15 efficacy of such services.

16 (g) DEFINITIONS.—In this section:

17 (1) ELIGIBLE INDIVIDUAL.—The term “eligible
18 individual” means, with respect to a State, an indi-
19 vidual who—

20 (A) is eligible for medical assistance under
21 the State plan under title XIX of the Social Se-
22 curity Act (or a waiver of such plan);

23 (B) is under the age of 21 (or, at the op-
24 tion of the State, under the age of 20, 19, or
25 18 as the State may choose), or in the case of

1 an individual described in section
2 1902(a)(10)(A)(i)(IX) of such Act (42 U.S.C.
3 1396a(a)(10)(A)(i)(IX)), under the age of 26;

4 (C) has been referred or admitted to an in-
5 tensive care unit, or has been seen by at least
6 one medical specialist, for a suspected genetic
7 or undiagnosed disease; or

8 (D) is suspected by at least one medical
9 specialist to have a neonatal-onset or pediatric-
10 onset genetic disease.

11 (2) GENETIC AND GENOMIC TESTING.—The
12 term “genetic and genomic testing”, with respect to
13 an eligible individual—

14 (A) means the determination of a sequence
15 of deoxyribonucleic acid bases in the genome of
16 such individual, and, if for the sole benefit of
17 the individual, a biological parent of such indi-
18 vidual for the purpose of determining whether
19 one or more potentially disease-causing genetic
20 variants are present in the genome of such indi-
21 vidual or such biological parent; and

22 (B) includes—

23 (i) the sequencing of the whole ge-
24 nome, the whole exome, or a panel of
25 genes; and

1 (ii) any analysis, interpretation, and
2 data report derived from such sequencing.

3 **SEC. 4. NATIONAL ACADEMY OF MEDICINE STUDY.**

4 (a) IN GENERAL.—Not later than one year after the
5 date of the enactment of this Act, the Secretary of Health
6 and Human Services shall enter into an arrangement with
7 the National Academy of Medicine under which the Acad-
8 emy agrees to study—

9 (1) how genetic and genomic testing may im-
10 prove preventative care and precision medicine;

11 (2) disparities in access to precision diagnostics
12 and associated therapeutics;

13 (3) how genetic and genomic testing may be
14 used to reduce health disparities in marginalized
15 communities;

16 (4) how the Federal Government may help to
17 reduce barriers to genetic and genomic testing, in-
18 cluding—

19 (A) encouraging the expansion of health
20 insurance coverage of genetic and genomic test-
21 ing, including diagnostic, predictive, and pre-
22 symptomatic testing, and genetic and genomic
23 testing (as defined in section 2(h)(2));

1 (B) supporting the collection of evidence
2 for the clinical utility and appropriate use of ge-
3 netic and genomic tests; and

4 (C) improving access to genetic counselors,
5 pathologists, and other relevant professions, in-
6 cluding strengthening related workforce edu-
7 cation and training efforts;

8 (5)(A) the extent to which coverage provisions
9 in the Medicare and Medicaid programs under titles
10 XVIII and XIX of the Social Security Act (42
11 U.S.C. 1395 et seq., 1396 et seq.) may restrain the
12 use of genetic and genomic testing that may improve
13 clinical outcomes for beneficiaries;

14 (B) the extent to which coverage provided pur-
15 suant to section 2 increased the use of genetic and
16 genomic testing and improved clinical outcomes for
17 beneficiaries; and

18 (C) how the Centers for Medicare & Medicaid
19 Services may make coverage determinations that
20 better suit a precision medicine approach to treat-
21 ment; and

22 (6) how genetic and genomic testing may im-
23 prove health outcomes for all pediatric populations
24 in the United States, including—

1 (A) children with a rare disease, including
2 a metabolic disease, neurologic disorder, or he-
3 reditary cancer testing in the presence of a sus-
4 pected or confirmed cancer diagnosis; and

5 (B) special populations, including—

6 (i) critically ill (non-infectious and
7 non-trauma) patients;

8 (ii) transplant patients;

9 (iii) individuals with cardiac disease;

10 and

11 (iv) individuals with, or who have a
12 family history of, a birth defect or develop-
13 mental disability.

14 (b) REPORT.—

15 (1) IN GENERAL.—The arrangement under sub-
16 section (a) shall provide for the National Academy
17 of Medicine to submit, not later than 2 years after
18 the date of enactment of this Act, a report on the
19 results of the study under subsection (a) to—

20 (A) the Secretary of Health and Human
21 Services;

22 (B) the Committee on Ways and Means
23 and the Committee on Energy and Commerce
24 of the House of Representatives; and

1 (C) the Committee on Finance and the
2 Committee on Health, Education, Labor, and
3 Pensions of the Senate.

4 (2) CONSULTATION.—The arrangement under
5 subsection (a) shall provide for the National Acad-
6 emy of Medicine, in developing the report required
7 by paragraph (1), to consult with physicians, other
8 health professionals, health educators, health profes-
9 sional organizations, relevant companies, patients,
10 patient organizations, the Health Resources and
11 Services Administration, the National Cancer Insti-
12 tute, the National Institutes of Health, the Agency
13 for Healthcare Research and Quality, and the Cen-
14 ters for Medicare & Medicaid Services.

15 (3) USE OF INFORMATION.—The National
16 Academy of Medicine shall, to the extent possible, in
17 conducting the study under subsection (a), utilize in-
18 formation included in the reports submitted pursu-
19 ant to subsections (f) and (g) of section 2.

20 **SEC. 5. CENTERS FOR MEDICARE & MEDICAID SERVICES**
21 **REPORT ON MEDICAID COVERAGE FOR GE-**
22 **NETIC AND GENOMIC TESTING.**

23 Not later than one year after the date of the enact-
24 ment of this Act, and annually thereafter for the subse-
25 quent 3 years, the Centers for Medicare & Medicaid Serv-

1 ices shall submit to the Secretary of Health and Human
2 Services, the Committees on Ways and Means and on En-
3 ergy and Commerce of the House of Representatives, and
4 the Committees on Finance and Health, Education,
5 Labor, and Pensions of the Senate a report on the extent
6 to which each of the 50 States provide coverage under the
7 State plan under title XIX of the Social Security Act (or
8 waiver of such plan) of genetic and genomic testing (as
9 defined in section 2(f)(2)) (including whole exome, whole
10 genome, gene panels, single gene tests, Chromosomal
11 microarray analysis, Fluorescence in situ hybridization,
12 and other genetic and genomic tests), including informa-
13 tion on—

14 (1) how often genetic and genomic diagnostic
15 testing services are covered and reimbursed;

16 (2) the frequency of denials for coverage and
17 the rationale for denying coverage;

18 (3) an analysis of which genetic and genomic
19 diagnostic tests are being approved or denied;

20 (4) how often test genetic counseling is covered
21 pre- and post-genetic and genomic diagnostic test-
22 ing;

23 (5) the turn-around time for prior authorization
24 requests; and

- 1 (6) any barriers to coverage of genetic and
- 2 genomic testing services identified.

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