Union Calendar No. 142

114TH CONGRESS 1ST SESSION

H. R. 6

[Report No. 114-190, Part I]

To accelerate the discovery, development, and delivery of 21st century cures, and for other purposes.

IN THE HOUSE OF REPRESENTATIVES

May 19, 2015

Mr. Upton (for himself, Ms. Degette, Mr. Pitts, Mr. Pallone, and Mr. Gene Green of Texas) 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

July 7, 2015

Additional sponsors: Mr. Barton, Mr. Cramer, Mr. Bucshon, Mr. Bili-RAKIS, Mrs. BLACKBURN, Mrs. BROOKS of Indiana, Mr. BURGESS, Mrs. ELLMERS of North Carolina, Mr. GRIFFITH, Mr. GUTHRIE, Mr. LANCE, Mr. McKinley, Mrs. McMorris Rodgers, Mr. Mullin, Mr. Murphy of Pennsylvania, Mr. Shimkus, Mr. Walden, Mr. Whitfield, Mr. Ros-KAM, Mr. HANNA, Mr. McCaul, Mrs. Comstock, Mr. Harris, Mr. MARCHANT, Mr. YARMUTH, Ms. CASTOR of Florida, Mr. LOEBSACK, Ms. Schakowsky, Mr. Tonko, Ms. Moore, Mr. Veasey, Mrs. Dingell, Mr. Fattah, Mr. Schrader, Mr. Nolan, Ms. Eshoo, Mr. Welch, Mr. DAVID SCOTT of Georgia, Mr. PERLMUTTER, Mr. COURTNEY, Mr. COHEN, Mr. DESAULNIER, Mr. LONG, Mr. BUTTERFIELD, Ms. CLARKE of New York, Mr. MICHAEL F. DOYLE of Pennsylvania, Mr. RUSH, Mr. ENGEL, Mr. McNerney, Ms. Matsui, Mr. Flores, Mr. Johnson of Ohio, Mr. GIBSON, Mr. KENNEDY, Mr. BEN RAY LUJÁN of New Mexico, Mr. Walz, Mr. Castro of Texas, Mrs. Bustos, Ms. Frankel of Florida, Ms. Brownley of California, Mr. Costa, Mrs. Wagner, Mr. Rod-NEY DAVIS of Illinois, Mr. Scalise, Mr. Latta, Mr. Harper, Mr. Olson, Mr. Kinzinger of Illinois, Mr. Pompeo, Mr. Collins of New York, Mrs. Mimi Walters of California, Mr. Allen, Mr. Sarbanes, Ms. Lee, Mrs. Miller of Michigan, Mr. Higgins, Mr. Huffman, Mr.

KILDEE, Mr. QUIGLEY, Mr. TAKAI, Mr. HECK of Nevada, Mr. PIERLUISI, Mr. Benishek, Mrs. Walorski, Mr. Israel, Mr. Hultgren, Ms. KUSTER, Mr. YODER, Mr. DENT, Mr. CURBELO of Florida, Mrs. KIRK-PATRICK, Mr. Polis, Mr. O'Rourke, Mr. Hudson, Mr. Cárdenas, Mrs. CAPPS, Mr. Roe of Tennessee, Mr. Bishop of Michigan, Mr. Ross, Mr. PAYNE, Ms. NORTON, Mr. DOLD, Mr. TURNER, Mr. COFFMAN, Mr. SIRES, Mr. MEADOWS, Mr. COSTELLO of Pennsylvania, Mr. GUTIÉRREZ, Mr. Messer, Mr. Walberg, Mr. Salmon, Mr. Jenkins of West Virginia, Mr. Pocan, Mr. Peters, Mrs. Torres, Ms. Lofgren, Ms. Meng, Mr. Takano, Mr. Pittenger, Mr. Denham, Mr. Barletta, Mr. HARDY, Mr. RUPPERSBERGER, Mr. VARGAS, Mr. GARAMENDI, Mr. HECK of Washington, Mr. Bera, Mr. Luetkemeyer, Mr. Zeldin, Ms. Clark of Massachusetts, Mr. Bost, Mrs. Lawrence, Ms. Velázquez, Mr. Rogers of Alabama, Ms. Edwards, Mr. Young of Iowa, Mr. Katko, Mr. Moulton, Mr. Fortenberry, Mr. McHenry, Mr. Hinojosa, Mr. Cartwright, Mr. Joyce, Mr. Valadao, Mr. Ashford, MOOLENAAR, Mr. DONOVAN, Mr. WENSTRUP, Mr. ELLISON, Mr. HAS-TINGS, Mrs. CAROLYN B. MALONEY of New York, Mr. NADLER, Ms. Wasserman Schultz, Mr. Trott, Ms. Kaptur, Ms. Kelly of Illinois, Mr. Rangel, Ms. Fudge, Mr. McGovern, Mr. Reed, Mr. Huizenga of Michigan, Mr. Lamalfa, Mr. Crenshaw, Ms. Sinema, Ms. McCol-LUM, Mr. Webster of Florida, Ms. Bordallo, Mr. Gibbs, Mr. Brendan F. Boyle of Pennsylvania, Mr. Conaway, Mr. Knight, Mr. Womack, Mr. Desjarlais, Mr. Royce, Mr. Diaz-Balart, Ms. STEFANIK, Mr. ZINKE, Mr. CAPUANO, Ms. JACKSON LEE, Mr. TIBERI, Ms. Herrera Beutler, Mr. Van Hollen, Mr. Guinta, Ms. Ros-LEHTINEN, Mr. SIMPSON, Ms. BONAMICI, Mr. CICILLINE, Mr. LIPINSKI, Miss Rice of New York, Mr. Paulsen, Ms. Speier, Mr. Cook, Mr. Langevin, Mr. Stivers, Mr. Cleaver, Mr. Collins of Georgia, Mr. HUNTER, Ms. HAHN, Mr. RIGELL, Mr. CHABOT, Ms. DELBENE, Mr. LoBiondo, Ms. Roybal-Allard, Mr. Perry, Mr. Delaney, Mr. Lewis, Mr. Ted Lieu of California, Mr. Johnson of Georgia, Ms. Wil-SON of Florida, Mr. Keating, Ms. Brown of Florida, Mr. Kind, Mr. Marino, Mr. Brady of Pennsylvania, Mr. Farr, Mr. Calvert, Mr. Larson of Connecticut, Ms. Judy Chu of California, Mr. Hoyer, Ms. EDDIE BERNICE JOHNSON of Texas, Ms. Maxine Waters of California, Mr. Lowenthal, Mr. Ryan of Ohio, Mrs. Davis of California, Mr. Kil-MER, and Ms. DUCKWORTH

July 7, 2015

Reported from the Committee on Energy and Commerce with an amendment [Strike out all after the enacting clause and insert the part printed in italic]

July 7, 2015

The Committee on Ways and Means discharged; committed to the Committee of the Whole House on the State of the Union and ordered to be printed

[For text of introduced bill, see copy of bill as introduced on May $19,\,2015$]

A BILL

To accelerate the discovery, development, and delivery of 21st century cures, 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,
- 3 SECTION 1. SHORT TITLE; TABLE OF CONTENTS.
- 4 (a) Short Title.—This Act may be cited as the "21st
- 5 Century Cures Act".
- 6 (b) Table of Contents for
- 7 this Act is as follows:
 - Sec. 1. Short title; table of contents.

TITLE I—DISCOVERY

Subtitle A—National Institutes of Health Funding

- Sec. 1001. National Institutes of Health reauthorization.
- Sec. 1002. NIH Innovation Fund.

Subtitle B—National Institutes of Health Planning and Administration

- Sec. 1021. NIH research strategic plan.
- Sec. 1022. Increasing accountability at the National Institutes of Health.
- Sec. 1023. Reducing administrative burdens of researchers.
- Sec. 1024. Exemption for the National Institutes of Health from the Paperwork Reduction Act requirements.
- Sec. 1025. NIH travel.
- Sec. 1026. Other transactions authority.
- Sec. 1027. NCATS phase IIB restriction.
- Sec. 1028. High-risk, high-reward research.
- Sec. 1029. Sense of Congress on increased inclusion of underrepresented communities in clinical trials.

Subtitle C—Supporting Young Emerging Scientists

- Sec. 1041. Improvement of loan repayment programs of the National Institutes of Health.
- Sec. 1042. Report.

Subtitle D—Capstone Grant Program

- Sec. 1061. Capstone award.
- Subtitle E—Promoting Pediatric Research Through the National Institutes of Health
- Sec. 1081. National pediatric research network.
- Sec. 1082. Global pediatric clinical study network sense of Congress.
- Sec. 1083. Appropriate age groupings in clinical research.

Subtitle F—Advancement of the National Institutes of Health Research and Data Access

- Sec. 1101. Sharing of data generated through NIH-funded research.
- Sec. 1102. Standardization of data in Clinical Trial Registry Data Bank on eligibility for clinical trials.

Subtitle G—Facilitating Collaborative Research

- Sec. 1121. Clinical trial data system.
- Sec. 1122. National neurological diseases surveillance system.
- Sec. 1123. Data on natural history of diseases.
- Sec. 1124. Accessing, sharing, and using health data for research purposes.

Subtitle H—Council for 21st Century Cures

Sec. 1141. Council for 21st Century Cures.

TITLE II—DEVELOPMENT

Subtitle A—Patient-Focused Drug Development

Sec. 2001. Development and use of patient experience data to enhance structured risk-benefit assessment framework.

Subtitle B—Qualification and Use of Drug Development Tools

- Sec. 2021. Qualification of drug development tools.
- Sec. 2022. Accelerated approval development plan.

Subtitle C—FDA Advancement of Precision Medicine

Sec. 2041. Precision medicine guidance and other programs of Food and Drug Administration.

Subtitle D—Modern Trial Design and Evidence Development

- Sec. 2061. Broader application of Bayesian statistics and adaptive trial designs.
- Sec. 2062. Utilizing evidence from clinical experience.
- Sec. 2063. Streamlined data review program.

Subtitle E—Expediting Patient Access

- Sec. 2081. Sense of Congress.
- Sec. 2082. Expanded access policy.
- Sec. 2083. Finalizing draft guidance on expanded access.

Subtitle F—Facilitating Responsible Manufacturer Communications

- Sec. 2101. Facilitating dissemination of health care economic information.
- Sec. 2102. Facilitating responsible communication of scientific and medical developments.

Subtitle G—Antibiotic Drug Development

- Sec. 2121. Approval of certain drugs for use in a limited population of patients.
- Sec. 2122. Susceptibility test interpretive criteria for microorganisms.
- Sec. 2123. Encouraging the development and use of new antimicrobial drugs.

Subtitle H-Vaccine Access, Certainty, and Innovation

- Sec. 2141. Timely review of vaccines by the Advisory Committee on Immunization Practices.
- Sec. 2142. Review of processes and consistency of ACIP recommendations.
- Sec. 2143. Meetings between CDC and vaccine developers.

Subtitle I—Orphan Product Extensions Now; Incentives for Certain Products for Limited Populations

- Sec. 2151. Extension of exclusivity periods for a drug approved for a new indication for a rare disease or condition.
- Sec. 2152. Reauthorization of rare pediatric disease priority review voucher incentive program.

Subtitle J—Domestic Manufacturing and Export Efficiencies

- Sec. 2161. Grants for studying the process of continuous drug manufacturing.
- Sec. 2162. Re-exportation among members of the European Economic Area.

Subtitle K—Enhancing Combination Products Review

Sec. 2181. Enhancing combination products review.

Subtitle L—Priority Review for Breakthrough Devices

Sec. 2201. Priority review for breakthrough devices.

Subtitle M—Medical Device Regulatory Process Improvements

- Sec. 2221. Third-party quality system assessment.
- Sec. 2222. Valid scientific evidence.
- Sec. 2223. Training and oversight in least burdensome appropriate means concept.
- Sec. 2224. Recognition of standards.
- Sec. 2225. Easing regulatory burden with respect to certain class I and class II devices.
- Sec. 2226. Advisory committee process.
- Sec. 2227. Humanitarian device exemption application.
- Sec. 2228. CLIA waiver study design guidance for in vitro diagnostics.

Subtitle N—Sensible Oversight for Technology Which Advances Regulatory Efficiency

- Sec. 2241. Health software.
- Sec. 2242. Applicability and inapplicability of regulation.
- Sec. 2243. Exclusion from definition of device.

Subtitle O—Streamlining Clinical Trials

- Sec. 2261. Protection of human subjects in research; applicability of rules.
- Sec. 2262. Use of non-local institutional review boards for review of investigational device exemptions and human device exemptions.
- Sec. 2263. Alteration or waiver of informed consent for clinical investigations.

Subtitle P—Improving Scientific Expertise and Outreach at FDA

- Sec. 2281. Silvio O. Conte Senior Biomedical Research Service.
- Sec. 2282. Enabling FDA scientific engagement.

- Sec. 2283. Reagan-Udall Foundation for the Food and Drug Administration.
- Sec. 2284. Collection of certain voluntary information exempted from Paperwork Reduction Act.
- Sec. 2285. Hiring authority for scientific, technical, and professional personnel.

Subtitle Q—Exempting From Sequestration Certain User Fees

Sec. 2301. Exempting from sequestration certain user fees of Food and Drug Administration.

TITLE III—DELIVERY

Subtitle A—Interoperability

Sec. 3001. Ensuring interoperability of health information technology.

Subtitle B—Telehealth

Sec. 3021. Telehealth services under the Medicare program.

Subtitle C—Encouraging Continuing Medical Education for Physicians

Sec. 3041. Exempting from manufacturer transparency reporting certain transfers used for educational purposes.

Subtitle D—Disposable Medical Technologies

Sec. 3061. Treatment of certain items and devices.

Subtitle E—Local Coverage Decision Reforms

Sec. 3081. Improvements in the Medicare local coverage determination (LCD) process.

Subtitle F—Medicare Pharmaceutical and Technology Ombudsman

Sec. 3101. Medicare pharmaceutical and technology ombudsman.

Subtitle G—Medicare Site-of-Service Price Transparency

Sec. 3121. Medicare site-of-Service price transparency.

Subtitle H—Medicare Part D Patient Safety and Drug Abuse Prevention

Sec. 3141. Programs to prevent prescription drug abuse under Medicare parts C and D.

TITLE IV—MEDICAID, MEDICARE, AND OTHER REFORMS

Subtitle A—Medicaid and Medicare Reforms

- Sec. 4001. Limiting Federal Medicaid reimbursement to States for durable medical equipment (DME) to Medicare payment rates.
- Sec. 4002. Medicare payment incentive for the transition from traditional x-ray imaging to digital radiography and other Medicare imaging payment provision.
- Sec. 4003. Implementation of Office of Inspector General recommendation to delay certain Medicare prescription drug plan prepayments.

Subtitle B—Cures Innovation Fund

Sec. 4041. Cures Innovation Fund.

Subtitle C—Other Reforms

Sec. 4061. SPR drawdown.

Subtitle D—Miscellaneous

Sec. 4081. Lyme disease and other tick-borne diseases.

TITLE I—DISCOVERY 1 Subtitle A—National Institutes of 2 Health Funding 3 SEC. 1001. NATIONAL INSTITUTES OF HEALTH REAUTHOR-5 IZATION. 6 Section 402A(a)(1) of the Public Health Service Act (42 U.S.C. 282a(a)(1)) is amended— 8 (1) in subparagraph (B), by striking at the end "and": 9 10 (2) in subparagraph (C), by striking at the end 11 the period and inserting a semicolon; and (3) by adding at the end the following new sub-12 13 paragraphs: 14 "(D) \$31,811,000,000 for fiscal year 2016; "(E) \$33,331,000,000 for fiscal year 2017; 15 16 and 17 "(F) \$34,851,000,000 for fiscal year 2018.". 18 SEC. 1002. NIH INNOVATION FUND. 19 (a) USE OF INNOVATION FUND.—Section 402(b) of the 20 Public Health Service Act (42 U.S.C. 282(b)) is amended—

1	(1) in paragraph (23), by striking at the end
2	"and";
3	(2) in paragraph (24), by striking at the end the
4	period and inserting "; and"; and
5	(3) by inserting after paragraph (24), the fol-
6	lowing new paragraph:
7	"(25) shall, with respect to funds appropriated
8	under section 402A(e) to the NIH Innovation Fund,
9	allocate such funds to the national research institutes
10	and national centers for conducting and supporting
11	innovation fund initiatives identified under para-
12	graph (3) of such section.".
13	(b) Establishment of Innovation Fund.—Section
14	402A of the Public Health Service Act (42 U.S.C. 282a)is
15	amended—
16	(1) by redesignating subsection (e) as subsection
17	(f); and
18	(2) by inserting after subsection (d) the following
19	new subsection:
20	"(e) NIH Innovation Fund.—
21	"(1) Establishment.—For the purpose of allo-
22	cations under section 402(b)(25), there is established
23	a fund to be known as the NIH Innovation Fund. The
24	Director of NIH shall, with respect to funds appro-
25	priated to the NIH Innovation Fund, allocate such

1	funds to support biomedical research through the
2	funding of basic, translational, and clinical research.
3	"(2) Amounts made available to fund.—
4	"(A) In general.—Subject to subpara-
5	graph (B), there is authorized to be appro-
6	priated, and appropriated, to the NIH Innova-
7	tion Fund out of any funds in the Treasury not
8	otherwise appropriated, \$2,000,000,000 for each
9	of fiscal years 2016 through 2020. The amounts
10	appropriated to the Fund by the preceding sen-
11	tence shall be in addition to any amounts other-
12	wise made available to the National Institutes of
13	Health.
14	"(B) Availability subject to appro-
15	PRIATIONS.—Amounts in the Fund shall not be
16	available except to the extent and in such
17	amounts as are provided in advance in appro-
18	priation Acts.
19	"(C) Allocation of amounts.—Of the
20	amounts made available from the NIH Innova-
21	tion Fund for allocations under section
22	402(b)(25) for a fiscal year—
23	"(i) not less than \$500,000,000 shall be
24	for the Accelerating Advancement Program
25	under paragraph (5);

1	"(ii) not less than 35 percent of such
2	amounts remaining after subtracting the al-
3	location for the Accelerating Advancement
4	Program shall be for early stage investiga-
5	tors (as defined in paragraph (7));
6	"(iii) not less than 20 percent of such
7	amounts remaining after subtracting the al-
8	location for the Accelerating Advancement
9	Program shall be for high-risk, high-reward
10	research under section 409K; and
11	"(iv) not more than 10 percent of such
12	amounts (without subtracting the allocation
13	for the Accelerating Advancement Program)
14	shall be for intramural research.
15	"(D) Inapplicability of certain provi-
16	SIONS.—Amounts in the NIH Innovation Fund
17	shall not be subject to—
18	"(i) any transfer authority of the Sec-
19	retary or the Director of NIH under section
20	241, subsection (c), subsection (d), or any
21	other provision of law (other than section
22	402(b)(25) and this subsection); or
23	"(ii) the Nonrecurring expenses fund
24	under section 223 of division G of the Con-

1	solidated Appropriations Act, 2008 (42
2	U.S.C. 3514a).
3	"(3) Authorized uses.—Amounts in the NIH
4	Innovation Fund established under paragraph (1)
5	may be used only to conduct or support innovative
6	biomedical research through the following:
7	"(A) Research in which—
8	"(i) a principal investigator has a spe-
9	cific project or specific objectives; and
10	"(ii) funding is tied to pursuit of such
11	project or objectives.
12	"(B) Research in which—
13	"(i) a principal investigator has shown
14	promise in biomedical research; and
15	"(ii) funding is not tied to a specific
16	project or specific objectives.
17	"(C) Research to be carried out by an early
18	stage investigator (as defined in paragraph (7)).
19	"(D) Research to be carried out by a small
20	business concern (as defined in section 3 of the
21	$Small\ Business\ Act).$
22	"(E) The Accelerating Advancement Pro-
23	gram under paragraph (5).
24	"(F) Development and implementation of
25	the strategic plan under paragraph (6).

1	"(4) Coordination.—In funding programs and
2	activities through the NIH Innovation Fund, the Sec-
3	retary, acting through the Director of NIH, shall—
4	"(A) ensure coordination among the na-
5	tional research institutes, the national centers,
6	and other departments, agencies, and offices of
7	the Federal Government; and
8	"(B) minimize unnecessary duplication.
9	"(5) Accelerating advancement program.—
10	The Director of NIH shall establish a program, to be
11	known as the Accelerating Advancement Program,
12	under which—
13	"(A) the Director of NIH partners with na-
14	tional research institutes and national centers to
15	accomplish important biomedical research objec-
16	tives; and
17	"(B) for every \$1 made available by the Di-
18	rector of NIH to a national research institute or
19	national center for a research project, the insti-
20	tute or center makes \$1 available for such project
21	from funds that are not derived from the NIH
22	$Innovation \ Fund.$
23	"(6) Strategic plan.—
24	"(A) In General.—The Director of NIH
25	shall ensure that scientifically based strategic

1	planning is implemented in support of research
2	priorities, including through development, use,
3	and updating of a research strategic plan that—
4	"(i) is designed to increase the efficient
5	and effective focus of biomedical research in
6	a manner that leverages the best scientific
7	opportunities through a deliberative plan-
8	ning process;
9	"(ii) identifies areas, to be known as
10	strategic focus areas, in which the resources
11	of the NIH Innovation Fund can contribute
12	to the goals of expanding knowledge to ad-
13	dress, and find more effective treatments for,
14	unmet medical needs in the United States,
15	including the areas of—
16	$``(I)\ biomarkers;$
17	"(II) precision medicine;
18	"(III) infectious diseases, includ-
19	ing pathogens listed as a qualifying
20	pathogen under section 505 $E(f)$ of the
21	Federal Food, Drug, and Cosmetic Act
22	or listed or designated as a tropical
23	disease under section 524 of such Act;
24	and
25	"(IV) antibiotics;

1	"(iii) includes objectives for each such
2	strategic focus area; and
3	"(iv) ensures that basic research re-
4	mains a priority.
5	"(B) UPDATES AND REVIEWS.—The Direc-
6	tor shall review and, as appropriate, update the
7	research strategic plan under subparagraph (A)
8	not less than every 18 months.
9	"(7) Definition.—In this subsection, the term
10	'early stage investigator' means an investigator
11	who—
12	"(A) will be the principal investigator or
13	the program director of the proposed research;
14	"(B) has never been awarded, or has been
15	awarded only once, a substantial, competing
16	grant by the National Institutes of Health for
17	independent research; and
18	"(C) is within 10 years of having com-
19	pleted—
20	"(i) the investigator's terminal degree;
21	or
22	"(ii) a medical residency (or the equiv-
23	alent).".
24	(c) Supplement, Not Supplant; Prohibition
25	AGAINST TRANSFER.—Funds appropriated pursuant to sec-

1	tion 402A(e) of the Public Health Service Act, as inserted
2	by subsection (b)—
3	(1) shall be used to supplement, not supplant, the
4	funds otherwise allocated by the National Institutes of
5	Health for biomedical research; and
6	(2) notwithstanding any transfer authority in
7	any appropriation Act, shall not be used for any pur-
8	pose other than allocating funds for conducting and
9	supporting innovation fund initiatives as described in
10	section 402(b)(25) of the Public Health Service Act,
11	as added by subsection (a).
12	Subtitle B—National Institutes of
13	Health Planning and Adminis-
14	tration
15	SEC. 1021. NIH RESEARCH STRATEGIC PLAN.
16	Section 402 of the Public Health Service Act (42
17	U.S.C. 282) is amended—
18	(1) in subsection (b), by amending paragraph
19	(5) to read as follows:
20	"(5) shall ensure that scientifically based stra-
21	tegic planning is implemented in support of research
22	priorities as determined by the agencies of the Na-
23	tional Institutes of Health, including through develop-
24	ment, use, and updating of the research strategic plan
25	under subsection (m);"; and

1	(2) by adding at the end the following:
2	"(m) Research Strategic Plan.—
3	"(1) Five-year plans for biomedical re-
4	SEARCH STRATEGY.—
5	"(A) In GENERAL.—For each successive
6	five-year period beginning with the period of fis-
7	cal years 2016 through 2020, the Director of
8	NIH, in consultation with the entities described
9	in subparagraph (B), shall develop and main-
10	tain a biomedical research strategic plan that—
11	"(i) is designed to increase the efficient
12	and effective focus of biomedical research in
13	a manner that leverages the best scientific
14	opportunities through a deliberative plan-
15	ning process;
16	"(ii) identifies areas, to be known as
17	strategic focus areas, in which the resources
18	of the National Institutes of Health can best
19	contribute to the goal of expanding knowl-
20	edge on human health in the United States
21	through biomedical research; and
22	"(iii) includes objectives for each such
23	strategic focus area.
24	"(B) Entities described.—The entities
25	described in this subparagraph are the directors

1	of the national research institutes and national
2	centers, researchers, patient advocacy groups,
3	and industry leaders.
4	"(2) Use of Plan.—The Director of NIH and
5	the directors of the national research institutes and
6	national centers shall use the strategic plan—
7	"(A) to identify research opportunities; and
8	"(B) to develop individual strategic plans
9	for the research activities of each of the national
10	research institutes and national centers that—
11	"(i) have a common template; and
12	"(ii) identify strategic focus areas in
13	which the resources of the national research
14	institutes and national centers can best con-
15	tribute to the goal of expanding knowledge
16	on human health in the United States
17	through biomedical research.
18	"(3) Contents of plans.—
19	"(A) Strategic focus areas.—The stra-
20	tegic focus areas identified pursuant to para-
21	$graph\ (1)(A)(ii)\ shall$ —
22	"(i) be identified in a manner that—
23	"(I) considers the return on in-
24	vestment to the United States public
25	through the investments of the National

1	Institutes of Health in biomedical re-
2	search; and
3	"(II) contributes to expanding
4	knowledge to improve the United
5	States public's health through bio-
6	medical research; and
7	"(ii) include overarching and trans-
8	National Institutes of Health strategic focus
9	areas, to be known as Mission Priority
10	Focus Areas, which best serve the goals of
11	preventing or eliminating the burden of a
12	disease or condition and scientifically merit
13	enhanced and focused research over the next
14	5 years.
15	"(B) Rare and pediatric diseases and
16	conditions.—In developing and maintaining a
17	strategic plan under this subsection, the Director
18	of NIH shall ensure that rare and pediatric dis-
19	eases and conditions remain a priority.
20	"(C) Workforce.—In developing and
21	maintaining a strategic plan under this sub-
22	section, the Director of NIH shall ensure that
23	maintaining the biomedical workforce of the fu-
24	ture, including the participation by scientists

1	from groups traditionally underrepresented in
2	the scientific workforce, remains a priority.
3	"(4) Initial plan.—Not later than 270 days
4	after the date of enactment of this subsection, the Di-
5	rector of NIH and the directors of the national re-
6	search institutes and national centers shall—
7	"(A) complete the initial strategic plan re-
8	quired by paragraphs (1) and (2); and
9	"(B) make such initial strategic plan pub-
10	licly available on the website of the National In-
11	stitutes of Health.
12	"(5) Review; updates.—
13	"(A) Progress reviews.—Not less than
14	annually, the Director of NIH, in consultation
15	with the directors of the national research insti-
16	tutes and national centers, shall conduct progress
17	reviews for each strategic focus area identified
18	$under\ paragraph\ (1)(A)(ii).$
19	"(B) UPDATES.—Not later than the end of
20	the 5-year period covered by the initial strategic
21	plan under this subsection, and every 5 years
22	thereafter, the Director of NIH, in consultation
23	with the directors of the national research insti-
24	tutes and national centers stakeholders in the

1	scientific field, advocates, and the public at
2	large, shall—
3	"(i) conduct a review of the plan, in-
4	cluding each strategic focus area identified
5	under paragraph (2)(B); and
6	"(ii) update such plan in accordance
7	with this section.".
8	SEC. 1022. INCREASING ACCOUNTABILITY AT THE NA-
9	TIONAL INSTITUTES OF HEALTH.
10	(a) Appointment and Terms of Directors of Na-
11	TIONAL RESEARCH INSTITUTES AND NATIONAL CEN-
12	TERS.—Subsection (a) of section 405 of the Public Health
13	Service Act (42 U.S.C. 284) is amended to read as follows:
14	"(a) Appointment; Terms.—
15	"(1) Appointment.—The Director of the Na-
16	tional Cancer Institute shall be appointed by the
17	President and the directors of the other national re-
18	search institutes, as well as the directors of the na-
19	tional centers, shall be appointed by the Director of
20	NIH. The directors of the national research institutes,
21	as well as national centers, shall report directly to the
22	Director of NIH.
23	"(2) TERMS.—

1	"(A) In general.—The term of office of a
2	director of a national research institute or na-
3	tional center shall be 5 years.
4	"(B) Removal.—The director of a national
5	research institute or national center may be re-
6	moved from office by the Director of NIH prior
7	to the expiration of such director's 5-year term.
8	"(C) Reappointment.—At the end of the
9	term of a director of a national research insti-
10	tute or national center, the director may be re-
11	appointed. There is no limit on the number of
12	terms a director may serve.
13	"(D) VACANCIES.—If the office of a director
14	of a national research institute or national cen-
15	ter becomes vacant before the end of such direc-
16	tor's term, the director appointed to fill the va-
17	cancy shall be appointed for a 5-year term start-
18	ing on the date of such appointment.
19	"(E) Transitional provision.—Each di-
20	rector of a national research institute or na-
21	tional center serving on the date of enactment of
22	the 21st Century Cures Act is deemed to be ap-
23	pointed for a 5-year term under this subsection

 $starting\ on\ such\ date\ of\ enactment.".$

24

1	(b) Compensation to Consultants or Individual
2	Scientists.—Section 202 of the Departments of Labor,
3	Health and Human Services, and Education, and Related
4	Agencies Appropriations Act, 1993 (Public Law 102–394;
5	42 U.S.C. 238f note) is amended by striking "portable
6	structures;" and all that follows and inserting "portable
7	structures.".
8	(c) Review of Certain Awards by Directors.—
9	Section 405(b) of the Public Health Service Act (42 U.S.C.
10	284(b)) is amended by adding at the end the following:
11	"(3) Before an award is made by a national research
12	institute or by a national center for a grant for a research
13	program or project (commonly referred to as an 'R-series
14	grant'), other than an award constituting a noncompeting
15	renewal of such grant, or a noncompeting administrative
16	supplement to such grant, the director of such national re-
17	search institute or national center—
18	"(A) shall review and approve the award; and
19	"(B) shall take into consideration—
20	"(i) the mission of the national research in-
21	stitute or national center and the scientific pri-
22	orities identified in the strategic plan under sec-
23	$tion \ 402(m); \ and$

1	"(ii) whether other agencies are funding
2	programs or projects to accomplish the same
3	goal.".
4	(d) IOM STUDY ON DUPLICATION IN FEDERAL BIO-
5	MEDICAL RESEARCH.—The Secretary of Health and
6	Human Services shall enter into an arrangement with the
7	Institute of Medicine of the National Academies (or, if the
8	Institute declines, another appropriate entity) under which
9	the Institute (or other appropriate entity) not later than
10	2 years after the date of enactment of this Act will—
11	(1) complete a study on the extent to which bio-
12	medical research conducted or supported by Federal
13	agencies is duplicative; and
14	(2) submit a report to the Congress on the results
15	of such study, including recommendations on how to
16	prevent such duplication.
17	SEC. 1023. REDUCING ADMINISTRATIVE BURDENS OF RE-
18	SEARCHERS.
19	(a) Plan Preparation and Implementation of
20	Measures To Reduce Administrative Burdens.—The
21	Director of the National Institutes of Health shall prepare
22	a plan, including time frames, and implement measures to
23	reduce the administrative burdens of researchers funded by
24	the National Institutes of Health, taking into account the

1	recommendations, evaluations, and plans researched by the
2	following entities:
3	(1) The Scientific Management Review Board.
4	(2) The National Academy of Sciences.
5	(3) The 2007 and 2012 Faculty Burden Survey
6	conducted by The Federal Demonstration Partner-
7	ship.
8	(4) Relevant recommendations from the Research
9	Business Models Working Group.
10	(b) Report.—Not later than two years after the date
11	of enactment of this Act, the Director of the National Insti-
12	tutes of Health shall submit to Congress a report on the
13	extent to which the Director has implemented measures pur-
14	suant to subsection (a).
15	SEC. 1024. EXEMPTION FOR THE NATIONAL INSTITUTES OF
16	HEALTH FROM THE PAPERWORK REDUCTION
17	ACT REQUIREMENTS.
18	Section 3518(c)(1) of title 44, United States Code, is
19	amended—
20	(1) in subparagraph (C), by striking "; or" and
21	inserting a semicolon;
22	(2) in subparagraph (D), by striking the period
23	at the end and inserting "; or"; and
24	(3) by inserting at the end the following new
25	subparagraph:

1	"(E) during the conduct of research by the Na-
2	tional Institutes of Health.".
3	SEC. 1025. NIH TRAVEL.
4	It is the sense of Congress that participation in or
5	sponsorship of scientific conferences and meetings is essen-
6	tial to the mission of the National Institutes of Health.
7	SEC. 1026. OTHER TRANSACTIONS AUTHORITY.
8	Section 480 of the Public Health Service Act (42
9	U.S.C. 287a) is amended—
10	(1) in subsection (b), by striking "the appropria-
11	tion of funds as described in subsection (g)" and in-
12	serting "the availability of funds as described in sub-
13	section (f)";
14	(2) in subsection (e)(3), by amending subpara-
15	graph (C) to read as follows:
16	"(C) Other transactions authority.—
17	The Director of the Center shall have other trans-
18	actions authority in entering into transactions
19	to fund projects in accordance with the terms
20	and conditions of this section.";
21	(3) by striking subsection (f); and
22	(4) by redesignating subsection (g) as subsection
23	<i>(f)</i> .

1	SEC. 1027. NCATS PHASE IIB RESTRICTION.
2	Section 479 of the Public Health Service Act (42
3	U.S.C. 287) is amended—
4	(1) prior to making the amendments under
5	paragraph (2), by striking "IIB" each place it ap-
6	pears and inserting "III"; and
7	(2) by striking "IIA" each place it appears and
8	inserting "IIB".
9	SEC. 1028. HIGH-RISK, HIGH-REWARD RESEARCH.
10	Part B of title IV of the Public Health Service Act
11	(42 U.S.C. 284 et seq.) is amended by adding at the end
12	the following:
13	"SEC. 409K. HIGH-RISK, HIGH-REWARD RESEARCH PRO-
13 14	"SEC. 409K. HIGH-RISK, HIGH-REWARD RESEARCH PROGRAM.
14	GRAM.
14 15	GRAM. "The director of each national research institute shall,
14 15 16	GRAM. "The director of each national research institute shall, as appropriate—
14 15 16 17	GRAM. "The director of each national research institute shall, as appropriate— "(1) establish programs to conduct or support re-
14 15 16 17	GRAM. "The director of each national research institute shall, as appropriate— "(1) establish programs to conduct or support research projects that pursue innovative approaches to
114 115 116 117 118	GRAM. "The director of each national research institute shall, as appropriate— "(1) establish programs to conduct or support research projects that pursue innovative approaches to major contemporary challenges in biomedical research
114 115 116 117 118 119 220	GRAM. "The director of each national research institute shall, as appropriate— "(1) establish programs to conduct or support research projects that pursue innovative approaches to major contemporary challenges in biomedical research that involve inherent high risk, but have the potential
14 15 16 17 18 19 20 21	"The director of each national research institute shall, as appropriate— "(1) establish programs to conduct or support research projects that pursue innovative approaches to major contemporary challenges in biomedical research that involve inherent high risk, but have the potential to lead to breakthroughs; and

1	SEC. 1029. SENSE OF CONGRESS ON INCREASED INCLUSION
2	OF UNDERREPRESENTED COMMUNITIES IN
3	CLINICAL TRIALS.
4	It is the sense of Congress that the National Institute
5	on Minority Health and Health Disparities (NIMHD)
6	should include within its strategic plan ways to increase
7	representation of underrepresented communities in clinical
8	trials.
9	Subtitle C—Supporting Young
10	Emerging Scientists
11	SEC. 1041. IMPROVEMENT OF LOAN REPAYMENT PROGRAMS
12	OF THE NATIONAL INSTITUTES OF HEALTH.
13	(a) In General.—Part G of title IV of the Public
14	Health Service (42 U.S.C. 288 et seq.) is amended—
15	(1) by redesignating the second section 487F (42
16	U.S.C. 288-6; relating to pediatric research loan re-
17	payment program) as section 487G; and
18	(2) by inserting after section 487G, as so redesig-
19	nated, the following:
20	"SEC. 487H. LOAN REPAYMENT PROGRAM.
21	"(a) In General.—The Secretary shall establish a
22	program, based on workforce and scientific needs, of enter-
23	ing into contracts with qualified health professionals under
24	which such health professionals agree to engage in research
25	in consideration of the Federal Government agreeing to
26	pay, for each year of engaging in such research, not more

- 1 than \$50,000 of the principal and interest of the edu-
- 2 cational loans of such health professionals.
- 3 "(b) Adjustment for Inflation.—Beginning with
- 4 respect to fiscal year 2017, the Secretary may increase the
- 5 maximum amount specified in subsection (a) by an amount
- 6 that is determined by the Secretary, on an annual basis,
- 7 to reflect inflation.
- 8 "(c) Limitation.—The Secretary may not enter into
- 9 a contract with a health professional pursuant to subsection
- 10 (a) unless such professional has a substantial amount of
- 11 educational loans relative to income.
- 12 "(d) Applicability of Certain Provisions Re-
- 13 Garding Obligated Service.—Except to the extent in-
- 14 consistent with this section, the provisions of sections 338B,
- 15 338C, and 338E shall apply to the program established
- 16 under this section to the same extent and in the same man-
- 17 ner as such provisions apply to the National Health Service
- 18 Corps Loan Repayment Program established under section
- 19 *338B*.
- 20 "(e) Availability of Appropriations.—Amounts
- 21 appropriated for a fiscal year for contracts under subsection
- 22 (a) are authorized to remain available until the expiration
- 23 of the second fiscal year beginning after the fiscal year for
- 24 which the amounts were appropriated.".

1	(b) UPDATE OF OTHER LOAN REPAYMENT PRO-
2	GRAMS.—
3	(1) Section 464z-5(a) of the Public Health Serv-
4	ice Act (42 U.S.C.285t-2(a)) is amended—
5	(A) by striking "\$35,000" and inserting
6	"\$50,000"; and
7	(B) by adding at the end the following new
8	sentence: "Subsection (b) of section 487H shall
9	apply with respect to the maximum amount
10	specified in this subsection in the same manner
11	as it applies to the maximum amount specified
12	in subsection (a) of such section.".
13	(2) Section 487A(a) of such Act (42 U.S.C. 288–
14	1(a)) is amended—
15	(A) by striking "\$35,000" and inserting
16	"\$50,000"; and
17	(B) by adding at the end the following new
18	sentence: "Subsection (b) of section 487H shall
19	apply with respect to the maximum amount
20	specified in this subsection in the same manner
21	as it applies to the maximum amount specified
22	in subsection (a) of such section.".
23	(3) Section 487B(a) of such Act (42 U.S.C. 288–
24	2(a)) is amended—

1	(A) by striking "\$35,000" and inserting
2	"\$50,000"; and
3	(B) by adding at the end the following new
4	sentence: "Subsection (b) of section 487H shall
5	apply with respect to the maximum amount
6	specified in this subsection in the same manner
7	as it applies to the maximum amount specified
8	in such subsection (a) of such section.".
9	(4) Section 487C(a)(1) of such Act (42 U.S.C.
10	288–3(a)(1)) is amended—
11	(A) by striking "\$35,000" and inserting
12	"\$50,000"; and
13	(B) by adding at the end the following new
14	sentence: "Subsection (b) of section 487H shall
15	apply with respect to the maximum amount
16	specified in this paragraph in the same manner
17	as it applies to the maximum amount specified
18	in such subsection (a) of such section.".
19	(5) Section $487E(a)(1)$ of such Act (42 U.S.C.
20	288–5(a)(1)) is amended—
21	(A) by striking "\$35,000" and inserting
22	"\$50,000"; and
23	(B) by adding at the end the following new
24	sentence: "Subsection (b) of section 487H shall
25	apply with respect to the maximum amount

1	specified in this paragraph in the same manner
2	as it applies to the maximum amount specified
3	in such subsection (a) of such section.".
4	(6) Section 487F(a) of such Act (42 U.S.C. 288–
5	5a(a)), as added by section 205 of Public Law 106-
6	505, is amended—
7	(A) by striking "\$35,000" and inserting
8	"\$50,000"; and
9	(B) by adding at the end the following new
10	sentence: "Subsection (b) of section 487H shall
11	apply with respect to the maximum amount
12	specified in this subsection in the same manner
13	as it applies to the maximum amount specified
14	in such subsection (a) of such section.".
15	(7) Section 487G of such Act (42 U.S.C. 288-6,
16	as redesignated by section 1041(a)(1)), is further
17	amended—
18	(A) in subsection $(a)(1)$, by striking
19	"\$35,000" and inserting "\$50,000"; and
20	(B) in subsection (b), by adding at the end
21	the following new sentence: "Subsection (b) of
22	section 487H shall apply with respect to the
23	maximum amount specified in subsection (a)(1)
24	in the same manner as it applies to the max-

1	imum amount specified in such subsection (a) of
2	such section.".
3	SEC. 1042. REPORT.
4	Not later than 18 months after the date of the enact-
5	ment of this Act, the Director of the National Institutes of
6	Health shall submit to Congress a report on efforts of the
7	National Institutes of Health to attract, retain, and develop
8	emerging scientists.
9	Subtitle D—Capstone Grant
10	Program
11	SEC. 1061. CAPSTONE AWARD.
12	Part G of title IV of the Public Health Service Act
13	(42 U.S.C. 288 et seq.) is amended by adding at the end
14	the following:
15	"SEC. 490. CAPSTONE AWARD.
16	"(a) In General.—The Secretary may make awards
17	(each of which, hereafter in this section, referred to as a
18	'Capstone Award') to support outstanding scientists who
19	have been funded by the National Institutes of Health.
20	"(b) Purpose.—Capstone Awards shall be made to fa-
21	cilitate the successful transition or conclusion of research
22	programs, or for other purposes, as determined by the Direc-
23	tor of NIH, in consultation with the directors of the na-
24	tional research institutes and national centers.

1	"(c) Duration and Amount.—The duration and
2	amount of each Capstone Award shall be determined by the
3	Director of NIH in consultation with the directors of the
4	national research institutes and national centers.
5	"(d) Limitation.—Individuals who have received a
6	Capstone Award shall not be eligible to have principle in-
7	vestigator status on subsequent awards from the National
8	Institutes of Health.".
9	Subtitle E—Promoting Pediatric
10	Research Through the National
11	Institutes of Health
12	SEC. 1081. NATIONAL PEDIATRIC RESEARCH NETWORK.
13	Section $409D(d)$ of the Public Health Service Act (42)
14	U.S.C. 284h(d)) is amended—
15	(1) in paragraph (1)—
16	(A) by striking "in consultation with the
17	Director of the Eunice Kennedy Shriver Na-
18	tional Institute of Child Health and Human De-
19	velopment and in collaboration with other ap-
20	propriate national research institutes and na-
21	tional centers that carry out activities involving
22	pediatric research" and inserting "in collabora-
23	tion with the national research institutes and
24	national centers that carry out activities involv-
25	ing pediatric research";

1	(B) by striking subparagraph (B);
2	(C) by striking "may be comprised of, as
3	appropriate" and all that follows through "the
4	pediatric research consortia" and inserting
5	"may be comprised of, as appropriate, the pedi-
6	atric research consortia"; and
7	(D) by striking "; or" at the end and insert-
8	ing a period; and
9	(2) in paragraph (1), paragraph (2)(A), the first
10	sentence of paragraph $(2)(E)$, and paragraph (4) , by
11	striking "may" each place it appears and inserting
12	"shall".
13	SEC. 1082. GLOBAL PEDIATRIC CLINICAL STUDY NETWORK
	SEC. 1082. GLOBAL PEDIATRIC CLINICAL STUDY NETWORK SENSE OF CONGRESS.
14	
13 14 15 16	SENSE OF CONGRESS.
141516	SENSE OF CONGRESS. It is the sense of Congress that—
14 15	SENSE OF CONGRESS. It is the sense of Congress that— (1) the National Institutes of Health should en-
14 15 16 17	SENSE OF CONGRESS. It is the sense of Congress that— (1) the National Institutes of Health should encourage a global pediatric clinical study network
14 15 16 17 18	SENSE OF CONGRESS. It is the sense of Congress that— (1) the National Institutes of Health should encourage a global pediatric clinical study network through the allocation of grants, contracts, or coopera-
14 15 16 17 18	SENSE OF CONGRESS. It is the sense of Congress that— (1) the National Institutes of Health should encourage a global pediatric clinical study network through the allocation of grants, contracts, or cooperative agreements to supplement the salaries of new and
14 15 16 17 18 19 20	SENSE OF CONGRESS. It is the sense of Congress that— (1) the National Institutes of Health should encourage a global pediatric clinical study network through the allocation of grants, contracts, or cooperative agreements to supplement the salaries of new and early investigators who participate in the global pedi-
14 15 16 17 18 19 20 21	SENSE OF CONGRESS. It is the sense of Congress that— (1) the National Institutes of Health should encourage a global pediatric clinical study network through the allocation of grants, contracts, or cooperative agreements to supplement the salaries of new and early investigators who participate in the global pediatric clinical study network;

1	new and early investigators, to entities that partici-
2	pate in the global pediatric clinical study network;
3	(3) the Food and Drug Administration should
4	engage the European Medicines Agency and other for-
5	eign regulatory entities during the formation of the
6	global pediatric clinical study network to encourage
7	their participation; and
8	(4) once a global pediatric clinical study network
9	is established and becomes operational, the Food and
10	Drug Administration should continue to engage the
11	European Medicines Agency and other foreign regu-
12	latory entities to encourage and facilitate their par-
13	ticipation in the network with the goal of enhancing
14	the global reach of the network.
15	SEC. 1083. APPROPRIATE AGE GROUPINGS IN CLINICAL RE-
16	SEARCH.
17	(a) Input From Experts.—Not later than 180 days
18	after the date of enactment of this Act, the Director of the
19	National Institutes of Health shall convene a workshop of
20	experts on pediatrics and experts on geriatrics to provide
21	input on—
22	(1) appropriate age groupings to be included in
23	research studies involving human subjects; and

1	(2) acceptable scientific justifications for exclud-
2	ing participants from a range of age groups from
3	human subjects research studies.
4	(b) Guidelines.—Not later than 180 days after the
5	conclusion of the workshop under subsection (a), the Direc-
6	tor of the National Institutes of Health shall publish guide-
7	lines—
8	(1) addressing the consideration of age as an in-
9	clusion variable in research involving human subjects;
10	and
11	(2) identifying criteria for justifications for any
12	age-related exclusions in such research.
13	(c) Public Availability of Findings and Conclu-
14	Sions.—The Director of the National Institutes of Health
15	shall—
16	(1) make the findings and conclusions resulting
17	from the workshop under subsection (a) available to
18	the public on the website of the National Institutes of
19	Health; and
20	(2) not less than biennially, disclose to the public
21	on such website the number of children included in re-
22	search that is conducted or supported by the National
23	Institutes of Health, disaggregated by developmentally
24	appropriate age group, race, and gender.

1	Subtitle F—Advancement of the Na-
2	tional Institutes of Health Re-
3	search and Data Access
4	SEC. 1101. SHARING OF DATA GENERATED THROUGH NIH-
5	FUNDED RESEARCH.
6	Section 402 of the Public Health Service Act (42
7	U.S.C. 282) (as amended by section 1021(2)) is further
8	amended by adding at the end the following:
9	"(n) Sharing of Data Generated Through NIH-
10	FUNDED RESEARCH.—
11	"(1) Authority.—Subject to paragraph (2), the
12	Director of NIH may require recipients of the award
13	of an NIH grant or other financial support, provided
14	that the research is fully funded through such grant
15	or other support, to share scientific data generated
16	from research conducted through such support for re-
17	search purposes.
18	"(2) Limitation.—The Director of NIH shall
19	not require the sharing of data that is inconsistent
20	with applicable law and policy protecting—
21	"(A) privacy and confidentiality;
22	"(B) proprietary interests;
23	"(C) business confidential information;
24	"(D) intellectual property rights; and
25	"(E) other relevant rights"

1	SEC. 1102. STANDARDIZATION OF DATA IN CLINICAL TRIAL
2	REGISTRY DATA BANK ON ELIGIBILITY FOR
3	CLINICAL TRIALS.
4	(a) Standardization.—
5	(1) In General.—Section 402(j) of the Public
6	Health Service Act (42 U.S.C. 282(j)) is amended—
7	(A) by redesignating paragraph (7) as
8	paragraph (8); and
9	(B) by inserting after paragraph (6) the fol-
10	lowing:
11	"(7) Standardization.—The Director of NIH
12	shall—
13	"(A) ensure that the registry and results
14	data bank is easily used by the public;
15	"(B) ensure that entries in the registry and
16	results data bank are easily compared;
17	"(C) ensure that information required to be
18	submitted to the registry and results data bank,
19	including recruitment information under para-
20	graph (2)(A)(ii)(II), is submitted by persons and
21	posted by the Director of NIH in a standardized
22	format and includes at least—
23	"(i) the disease or indication being
24	studied;

1	"(ii) inclusion criteria such as age,
2	gender, diagnosis or diagnoses, laboratory
3	values, or imaging results; and
4	"(iii) exclusion criteria such as specific
5	diagnosis or diagnoses, laboratory values, or
6	prohibited medications; and
7	"(D) to the extent possible, in carrying out
8	this paragraph, make use of standard health care
9	terminologies, such as the International Classi-
10	fication of Diseases or the Current Procedural
11	Terminology, that facilitate electronic matching
12	to data in electronic health records or other rel-
13	evant health information technologies.".
14	(2) Conforming amendment.—Clause (iv) of
15	section 402(j)(2)(B) of the Public Health Service Act
16	$(42\ U.S.C.\ 282(j)(2)(B))$ is hereby stricken.
17	(b) Consultation.—Not later than 90 days after the
18	date of enactment of this Act, the Secretary of Health and
19	Human Services shall consult with stakeholders (including
20	patients, researchers, physicians, industry representatives,
21	health information technology providers, the Food and Drug
22	Administration, and standard setting organizations such as
23	CDISC that have experience working with Federal agencies
24	to standardize health data submissions) to receive advice
25	on enhancements to the clinical trial registry data bank

- 1 under section 402(j) of the Public Health Service Act (42
- 2 U.S.C. 282(j)) (including enhancements to usability,
- 3 functionality, and search capability) that are necessary to
- 4 implement paragraph (7) of section 402(j) of such Act, as
- 5 added by subsection (a).
- 6 (c) Applicability.—Not later than 18 months after
- 7 the date of enactment of this Act, the Secretary of Health
- 8 and Human Services shall begin implementation of para-
- 9 graph (7) of section 402(j) of the Public Health Service Act,
- 10 as added by subsection (a).

11 Subtitle G—Facilitating

12 Collaborative Research

- 13 SEC. 1121. CLINICAL TRIAL DATA SYSTEM.
- 14 (a) Establishment.—The Secretary, acting through
- 15 the Commissioner of Food and Drugs and the Director of
- 16 the National Institutes of Health, shall enter into a coopera-
- 17 tive agreement, contract, or grant for a period of 7 years,
- 18 to be known as the Clinical Trial Data System Agreement,
- 19 with one or more eligible entities to implement a pilot pro-
- 20 gram with respect to all clinical trial data obtained from
- 21 qualified clinical trials for purposes of registered users con-
- 22 ducting further research on such data.
- 23 (b) Application.—Eligible entities seeking to enter
- 24 into a cooperative agreement, contract, or grant with the
- 25 Secretary under this section shall submit to the Secretary

- 1 an application in such time and manner, and containing
- 2 such information, as the Secretary may require in accord-
- 3 ance with this section. The Secretary shall not enter into
- 4 a cooperative agreement, contract, or grant under this sec-
- 5 tion with an eligible entity unless such entity submits an
- 6 application including the following:
- 7 (1) A certification that the eligible entity is not 8 currently and does not plan to be involved in spon-9 soring, operating, or participating in a clinical trial 10 nor collaborating with another entity for the purposes 11 of sponsoring, operating, or participating in a clin-12 ical trial.
 - (2) Information demonstrating that the eligible entity can compile clinical trial data in standardized formats using terminologies and standards that have been developed by recognized standards developing organizations with input from diverse stakeholder groups, and information demonstrating that the eligible entity can de-identify clinical trial data consistent with the requirements of section 164.514 of title 45, Code of Federal Regulations (or successor regulations).
 - (3) A description of the system the eligible entity will use to store and maintain such data, and information demonstrating that this system will comply

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- with applicable standards and requirements for en suring the security of the clinical trial data.
 - (4) A certification that the eligible entity will allow only registered users to access and use de-identified clinical trial data, gathered from qualified clinical trials, and that the eligible entity will allow each registered user to access and use such data only after such registered user agrees in writing to the terms described in (e)(4)(B), and such other carefully controlled contractual terms as may be defined by the Secretary.
 - (5) Evidence demonstrating the ability of the eligible entity to ensure that registered users disseminate the results of the research conducted in accordance with this section to interested parties to serve as a guide to future medical product development or scientific research.
 - (6) The plan of the eligible entity for securing funding for the activities it would conduct under the clinical trial data system agreement from governmental sources and private foundations, entities, and individuals.
 - (7) Evidence demonstrating a proven track record of—

1	(A) being a neutral third party in working
2	with medical product manufacturers, academic
3	institutions, and the Food and Drug Adminis-
4	tration; and
5	(B) having the ability to protect confiden-
6	$tial\ data.$
7	(8) An agreement that the eligible entity will
8	work with the Comptroller General of the United
9	States for purposes of the study and report under sub-
10	section (d).
11	(c) Extension, Expansion, Termination.—The Sec-
12	retary, acting through the Commissioner of Food and Drugs
13	and the Director of the National Institutes of Health, upon
14	the expiration of the 7-year period referred to in subsection
15	(a), may extend (including permanently), expand, or termi-
16	nate the pilot program established under such subsection,
17	in whole or in part.
18	(d) Study and Report.—
19	(1) In General.—The Comptroller General of
20	the United States shall conduct a study and issue a
21	report to the Congress and the Secretary with respect
22	to the pilot program established under subsection (a),
23	not later than 6 years after the date on which the
24	pilot program is established under subsection (a).

1	(2) STUDY.—The study under paragraph (1)
2	shall—
3	(A) review the effectiveness of the pilot pro-
4	gram established under subsection (a); and
5	(B) be designed to formulate recommenda-
6	tions on improvements to the program.
7	(3) Report.—The report under paragraph (1)
8	shall contain at least the following information:
9	(A) The new discoveries, research inquiries,
10	or clinical trials that have resulted from access-
11	ing clinical trial data under the pilot program
12	established under subsection (a).
13	(B) The number of times scientists have
14	accessed such data, disaggregated by research
15	area and clinical trial phase.
16	(C) An analysis of whether the program has
17	helped to reduce adverse events in clinical trials.
18	(D) An analysis of whether scientists have
19	raised any concerns about the burden of having
20	to share data with the system established under
21	the program and a description, if any, of such
22	burden.
23	(E) An analysis of privacy and data integ-
24	rity practices used in the program.
25	(e) Definitions.—In this section:

1	(1) The term "eligible entity" means an entity
2	that has experienced personnel with clinical and other
3	technical expertise in the biomedical sciences and bio-
4	medical ethics and that is—
5	(A) an institution of higher education (as
6	such term is defined in section 1001 of the High-
7	er Education Act of 1965 (20 U.S.C. 1001)) or
8	a consortium of such institutions; or
9	(B) an organization described in section
10	501(c)(3) of title 26 of the Internal Revenue Code
11	of 1986 and exempt from tax under section
12	501(a) of such title.
13	(2) The term "medical product" means a drug
14	(as defined in section 201(g) of the Federal Food,
15	Drug, and Cosmetic Act (21 U.S.C. 331(g))), a device
16	(as defined in section 201(h) of such Act (21 U.S.C.
17	331(h)), a biological product (as defined in section
18	351 of the Public Health Service Act (42 U.S.C.
19	262)), or any combination thereof.
20	(3) The term "qualified clinical trial" means a
21	clinical trial sponsored solely by an agency of the De-
22	partment of Health and Human Services with respect
23	to a medical product—
24	(A) that—

1	(i) was approved or cleared under sec-
2	tion 505, 510(k), or 515, or has an exemp-
3	tion for investigational use in effect under
4	section 505 or 520(m), of the Federal Food,
5	Drug, and Cosmetic Act (42 U.S.C. 301 et
6	seq.); or
7	(ii) was licensed under section 351 of
8	the Public Health Service Act (42 U.S.C.
9	262) or has an exemption for investiga-
10	tional use in effect under such section 351;
11	or
12	(B) that is an investigational product for
13	which the original development was discontinued
14	and with respect to which—
15	(i) no additional work to support ap-
16	proval, licensure, or clearance of such med-
17	ical product is being or is planned to be un-
18	dertaken by the sponsor of the original de-
19	velopment program, its successors, assigns,
20	or collaborators; and
21	(ii) the sponsor of the original inves-
22	tigational development program has pro-
23	vided its consent to the Secretary for inclu-
24	sion of data regarding such product in the
25	sustem established under this section.

1	(4) The term "registered user" means a scientific
2	or medical researcher who has—
3	(A) a legitimate biomedical research pur-
4	pose for accessing information from the clinical
5	trials data system and has appropriate quali-
6	fications to conduct such research; and
7	(B) agreed in writing not to transfer to any
8	other person that is not a registered user de-iden-
9	tified clinical trial data from qualified clinical
10	trials accessed through an eligible entity, use
11	such data for reasons not specified in the re-
12	search proposal, or seek to re-identify qualified
13	clinical trial participants.
14	(5) The term "Secretary" means the Secretary of
15	Health and Human Services.
16	SEC. 1122. NATIONAL NEUROLOGICAL DISEASES SURVEIL-
17	LANCE SYSTEM.
18	Part P of title III of the Public Health Service Act
19	(42 U.S.C. 280g et seq.) is amended by adding at the end
20	the following:
21	"SEC. 399V-6 SURVEILLANCE OF NEUROLOGICAL DISEASES.
22	"(a) In General.—The Secretary, acting through the
23	Director of the Centers for Disease Control and Prevention
24	and in coordination with other agencies as determined ap-
25	propriate by the Secretary, shall—

1	"(1) enhance and expand infrastructure and ac-
2	tivities to track the epidemiology of neurological dis-
3	eases, including multiple sclerosis and Parkinson's
4	disease; and
5	"(2) incorporate information obtained through
6	such activities into a statistically sound, scientifically
7	credible, integrated surveillance system, to be known
8	as the National Neurological Diseases Surveillance
9	System.
10	"(b) Research.—The Secretary shall ensure that the
11	National Neurological Diseases Surveillance System is de-
12	signed in a manner that facilitates further research on neu-
13	rological diseases.
14	"(c) Content.—In carrying out subsection (a), the
15	Secretary—
16	"(1) shall provide for the collection and storage
17	of information on the incidence and prevalence of
18	neurological diseases in the United States;
19	"(2) to the extent practicable, shall provide for
20	the collection and storage of other available informa-
21	tion on neurological diseases, such as information
22	concerning—
23	"(A) demographics and other information
24	associated or possibly associated with neuro-

1	logical diseases, such as age, race, ethnicity, sex,
2	geographic location, and family history;
3	"(B) risk factors associated or possibly asso-
4	ciated with neurological diseases, including ge-
5	netic and environmental risk factors; and
6	"(C) diagnosis and progression markers;
7	"(3) may provide for the collection and storage
8	of information relevant to analysis on neurological
9	diseases, such as information concerning—
10	"(A) the epidemiology of the diseases;
11	"(B) the natural history of the diseases;
12	"(C) the prevention of the diseases;
13	"(D) the detection, management, and treat-
14	ment approaches for the diseases; and
15	$\lq\lq(E)$ the development of outcomes measures;
16	and
17	"(4) may address issues identified during the
18	consultation process under subsection (d).
19	"(d) Consultation.—In carrying out this section, the
20	Secretary shall consult with individuals with appropriate
21	expertise, including—
22	"(1) epidemiologists with experience in disease
23	surveillance or registries;
24	"(2) representatives of national voluntary health
25	associations that—

1	"(A) focus on neurological diseases, includ-
2	ing multiple sclerosis and Parkinson's disease;
3	and
4	"(B) have demonstrated experience in re-
5	search, care, or patient services;
6	"(3) health information technology experts or
7	$other\ information\ management\ specialists;$
8	"(4) clinicians with expertise in neurological
9	diseases; and
10	"(5) research scientists with experience con-
11	ducting translational research or utilizing surveil-
12	lance systems for scientific research purposes.
13	"(e) Grants.—The Secretary may award grants to,
14	or enter into contracts or cooperative agreements with, pub-
15	lic or private nonprofit entities to carry out activities under
16	this section.
17	"(f) Coordination With Other Federal, State,
18	AND LOCAL AGENCIES.—Subject to subsection (h), the Sec-
19	retary shall make information and analysis in the National
20	Neurological Diseases Surveillance System available, as ap-
21	propriate—
22	"(1) to Federal departments and agencies, such
23	as the National Institutes of Health, the Food and
24	Drug Administration, the Centers for Medicare &
25	Medicaid Services, the Agency for Healthcare Re-

1	search and Quality, the Department of Veterans Af-
2	fairs, and the Department of Defense; and
3	"(2) to State and local agencies.
4	"(g) Public Access.—Subject to subsection (h), the
5	Secretary shall make information and analysis in the Na-
6	tional Neurological Diseases Surveillance System available,
7	as appropriate, to the public, including researchers.
8	"(h) Privacy.—The Secretary shall ensure that pri-
9	vacy and security protections applicable to the National
10	Neurological Diseases Surveillance System are at least as
11	stringent as the privacy and security protections under
12	HIPAA privacy and security law (as defined in section
13	3009(a)(2)).
14	"(i) Report.—Not later than 4 years after the date
15	of the enactment of this section, the Secretary shall submit
16	a report to the Congress concerning the implementation of
17	this section. Such report shall include information on—
18	"(1) the development and maintenance of the
19	National Neurological Diseases Surveillance System;
20	"(2) the type of information collected and stored
21	in the System;
22	"(3) the use and availability of such informa-
23	tion including avidelines for such use: and

1	"(4) the use and coordination of databases that
2	collect or maintain information on neurological dis-
3	eases.
4	"(j) Definition.—In this section, the term 'national
5	voluntary health association' means a national nonprofit
6	organization with chapters, other affiliated organizations,
7	or networks in States throughout the United States.
8	"(k) Authorization of Appropriations.—To carry
9	out this section, there is authorized to be appropriated
10	\$5,000,000 for each of fiscal years 2016 through 2020.".
11	SEC. 1123. DATA ON NATURAL HISTORY OF DISEASES.
12	(a) Sense of Congress.—It is the sense of the Con-
13	gress that studies on the natural history of diseases can help
14	to facilitate and expedite the development of medical prod-
15	ucts for such diseases.
16	(b) Authority.—Part A of title II of the Public
17	Health Service Act (42 U.S.C. 202 et seq.) is amended by
18	adding at the end the following:
19	"SEC. 229A. DATA ON NATURAL HISTORY OF DISEASES.
20	"(a) In General.—The Secretary may, for the pur-
21	poses described in subsection (b)—
22	"(1) participate in public-private partnerships
23	engaged in one or more activities specified in sub-
24	section (c); and

1	"(2) award grants to patient advocacy groups or
2	other organizations determined appropriate by the
3	Secretary.
4	"(b) Purposes Described.—The purposes described
5	in this subsection are to establish or facilitate the collection,
6	maintenance, analysis, and interpretation of data regard-
7	ing the natural history of diseases, with a particular focus
8	on rare diseases.
9	"(c) Activities of Public-Private Partner-
10	SHIPS.—The activities of public-private partnerships in
11	which the Secretary may participate for purposes of this
12	section include—
13	"(1) cooperating with other entities that sponsor
14	or maintain disease registries, including disease reg-
15	istries and disease registry platforms for rare dis-
16	eases;
17	"(2) developing or enhancing a secure informa-
18	tion technology system that—
19	"(A) has the capacity to support data needs
20	across a wide range of disease studies;
21	"(B) is easily modified as knowledge is
22	gained during such studies; and
23	"(C) is capable of handling increasing
24	amounts of data as more studies are carried out;
25	and

1	"(3) providing advice to clinical researchers, pa-
2	tient advocacy groups, and other entities with respect
3	to—
4	"(A) the design and conduct of disease stud-
5	ies;
6	"(B) the modification of any such ongoing
7	studies; and
8	"(C) addressing associated patient privacy
9	issues.
10	"(d) Availability of Data on Natural History of
11	Diseases.—Data relating to the natural history of diseases
12	obtained, aggregated, or otherwise maintained by a public-
13	private partnership in which the Secretary participates
14	under subsection (a) shall be made available, consistent
15	with otherwise applicable Federal and State privacy laws,
16	to the public (including patient advocacy groups, research-
17	ers, and drug developers) to help to facilitate and expedite
18	medical product development programs.
19	$\hbox{\it ``(e)} Confidentiality. \hbox{\itNotwith standing } \ subsection$
20	(d), nothing in this section authorizes the disclosure of any
21	information that is a trade secret or commercial or finan-
22	cial information that is privileged or confidential and sub-
23	$ject\ to\ section\ 552(b)(4)\ of\ title\ 5,\ United\ States\ Code,\ or$
24	section 1905 of title 18, United States Code.

1	"(f) Authorization of Appropriations.—There is
2	authorized to be appropriated to carry out this section
3	\$5,000,000 for each of fiscal years 2016 through 2020.".
4	SEC. 1124. ACCESSING, SHARING, AND USING HEALTH DATA
5	FOR RESEARCH PURPOSES.
6	(a) In General.—(1) The HITECH Act (title XIII
7	of division A of Public Law 111-5) is amended by adding
8	at the end of subtitle D of such Act (42 U.S.C. 17921 et
9	seq.) the following:
10	"PART 4—ACCESSING, SHARING, AND USING
11	HEALTH DATA FOR RESEARCH PURPOSES
12	"SEC. 13441. REFERENCES.
13	"In this part:
14	"(1) The Rule.—References to 'the Rule' refer
15	to part 160 or part 164, as appropriate, of title 45,
16	Code of Federal Regulations (or any successor regula-
17	tion).
18	"(2) PART 164.—References to a specified section
19	of 'part 164', refer to such specified section of part
20	164 of title 45, Code of Federal Regulations (or any
21	$successor\ section).$
22	"SEC. 13442. DEFINING HEALTH DATA RESEARCH AS PART
23	OF HEALTH CARE OPERATIONS.
24	"(a) In General.—Subject to subsection (b), the Sec-
25	retary shall revise or clarify the Rule to allow the use and

1	disclosure of protected health information by a covered enti-
2	ty for research purposes, including studies whose purpose
3	is to obtain generalizable knowledge, to be treated as the
4	use and disclosure of such information for health care oper-
5	ations described in subparagraph (1) of the definition of
6	health care operations in section 164.501 of part 164.
7	"(b) Modifications to Rules for Disclosures
8	FOR HEALTH CARE OPERATIONS.—In applying section
9	164.506 of part 164 to the disclosure of protected health in-
10	formation described in subsection (a)—
11	"(1) the Secretary shall revise or clarify the Rule
12	so that the disclosure may be made by the covered en-
13	tity to only—
14	"(A) another covered entity for health care
15	operations (as defined in section 164.501 of part
16	164);
17	"(B) a business associate that has entered
18	into a contract under section 164.504(e) of part
19	164 with a disclosing covered entity to perform
20	health care operations; or
21	"(C) a business associate that has entered
22	into a contract under section 164.504(e) of part
23	164 for the purpose of data aggregation (as de-
24	fined in section 164,501 of part 164); and

1	"(2) the Secretary shall further revise or clarify
2	the Rule so that the limitation specified by section
3	164.506(c)(4) of part 164 does not apply to disclo-
4	sures that are described by subsection (a).
5	"(c) Rule of Construction.—This section shall not
6	be construed as prohibiting or restricting a use or disclosure
7	of protected health information for research purposes that
8	is otherwise permitted under part 164.
9	"SEC. 13443. TREATING DISCLOSURES OF PROTECTED
10	HEALTH INFORMATION FOR RESEARCH SIMI
11	LARLY TO DISCLOSURES OF SUCH INFORMA
	LARLY TO DISCLOSURES OF SUCH INFORMA TION FOR PUBLIC HEALTH PURPOSES.
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12 13	TION FOR PUBLIC HEALTH PURPOSES.
12 13 14	TION FOR PUBLIC HEALTH PURPOSES. "(a) REMUNERATION.—The Secretary shall revise on
12 13 14 15	TION FOR PUBLIC HEALTH PURPOSES. "(a) REMUNERATION.—The Secretary shall revise of clarify the Rule so that disclosures of protected health information.
12 13 14 15	TION FOR PUBLIC HEALTH PURPOSES. "(a) REMUNERATION.—The Secretary shall revise of clarify the Rule so that disclosures of protected health information for research purposes are not subject to the limitary
12 13 14 15 16	"(a) Remuneration.—The Secretary shall revise of clarify the Rule so that disclosures of protected health information for research purposes are not subject to the limitation on remuneration described in section
12 13 14 15 16 17	TION FOR PUBLIC HEALTH PURPOSES. "(a) REMUNERATION.—The Secretary shall revise of clarify the Rule so that disclosures of protected health information for research purposes are not subject to the limitation on remuneration described in section $164.502(a)(5)(ii)(B)(2)(ii)$ of part $164.$
12 13 14 15 16 17 18	"(a) Remuneration.—The Secretary shall revise of clarify the Rule so that disclosures of protected health information for research purposes are not subject to the limitation on remuneration described in section 164.502(a)(5)(ii)(B)(2)(ii) of part 164. "(b) Permitted Uses and Disclosures.—The Secretary
12 13 14 15 16 17 18 19	"(a) Remuneration.—The Secretary shall revise of clarify the Rule so that disclosures of protected health information for research purposes are not subject to the limitation on remuneration described in section 164.502(a)(5)(ii)(B)(2)(ii) of part 164. "(b) Permitted Uses and Disclosures.—The Secretary shall revise or clarify the Rule so that research acceptable.
18 19 20 21	"(a) Remuneration.—The Secretary shall revise of clarify the Rule so that disclosures of protected health information for research purposes are not subject to the limitation on remuneration described in section 164.502(a)(5)(ii)(B)(2)(ii) of part 164. "(b) Permitted Uses and Disclosures.—The Secretary shall revise or clarify the Rule so that research activities, including comparative research activities, related

23 are included as public health activities for purposes of

24 which a covered entity may disclose protected health infor-

1	mation to a person described in section 164.512(b)(1)(iii)
2	of part 164.
3	"SEC. 13444. PERMITTING REMOTE ACCESS TO PROTECTED
4	HEALTH INFORMATION BY RESEARCHERS.
5	"The Secretary shall revise or clarify the Rule so that
6	subparagraph (B) of section 164.512(i)(1)(ii) of part 164
7	(prohibiting the removal of protected health information by
8	a researcher) shall not prohibit remote access to health in-
9	formation by a researcher so long as—
10	"(1) appropriate security and privacy safe-
11	guards are maintained by the covered entity and the
12	researcher; and
13	"(2) the protected health information is not cop-
14	ied or otherwise retained by the researcher.
15	"SEC. 13445. ALLOWING ONE-TIME AUTHORIZATION OF USE
16	AND DISCLOSURE OF PROTECTED HEALTH
17	INFORMATION FOR RESEARCH PURPOSES.
18	"(a) In General.—The Secretary shall revise or clar-
19	ify the Rule to specify that an authorization for the use
20	or disclosure of protected health information, with respect
21	to an individual, for future research purposes shall be
22	deemed to contain a sufficient description of the purpose
23	of the use or disclosure if the authorization—
24	"(1) sufficiently describes the purposes such that
25	it would be reasonable for the individual to expect

1	that the protected health information could be used or
2	disclosed for such future research;
3	"(2) either—
4	"(A) states that the authorization will ex-
5	pire on a particular date or on the occurrence of
6	a particular event; or
7	"(B) states that the authorization will re-
8	main valid unless and until it is revoked by the
9	individual; and
10	"(3) provides instruction to the individual on
11	how to revoke such authorization at any time.
12	"(b) Revocation of Authorization.—The Sec-
13	retary shall revise or clarify the Rule to specify that, if an
14	individual revokes an authorization for future research pur-
15	poses such as is described by subsection (a), the covered enti-
16	ty may not make any further uses or disclosures based on
17	that authorization, except, as provided in paragraph (b)(5)
18	of section 164.508 of part 164, to the extent that the covered
19	entity has taken action in reliance on the authorization.".
20	(2) The table of sections in section 13001(b) of such
21	Act is amended by adding at the end of the items relating
22	to subtitle D the following new items:
	"Part 4—Accessing, Sharing, and Using Health Data for Research Purposes

[&]quot;Sec. 13441. References.

[&]quot;Sec. 13442. Defining health data research as part of health care operations.

[&]quot;Sec. 13443. Treating disclosures of protected health information for research similarly to disclosures of such information for public health purposes.

- "Sec. 13444. Permitting remote access to protected health information by researchers.
- "Sec. 13445. Allowing one-time authorization of use and disclosure of protected health information for research purposes.".
- 1 (b) Revision of Regulations.—Not later than 12
- 2 months after the date of the enactment of this Act, the Sec-
- 3 retary of Health and Human Services shall revise and clar-
- 4 ify the provisions of title 45, Code of Federal Regulations,
- 5 for consistency with part 4 of subtitle D of the HITECH
- 6 Act, as added by subsection (a).

7 Subtitle H—Council for 21st

8 Century Cures

- 9 SEC. 1141. COUNCIL FOR 21ST CENTURY CURES.
- 10 Title II of the Public Health Service Act (42 U.S.C.
- 11 202 et seq.) is amended by adding at the end the following:
- 12 "PART E—COUNCIL FOR 21ST CENTURY CURES
- 13 "SEC. 281. ESTABLISHMENT.
- 14 "A nonprofit corporation to be known as the Council
- 15 for 21st Century Cures (referred to in this part as the
- 16 'Council') shall be established in accordance with this sec-
- 17 tion. The Council shall be a public-private partnership
- 18 headed by an Executive Director (referred to in this part
- 19 as the 'Executive Director'), appointed by the members of
- 20 the Board of Directors. The Council shall not be an agency
- 21 or instrumentality of the United States Government.

"SEC. 281A. PURPOSE.

2	"The purpose of the Council is to accelerate the dis-
3	covery, development, and delivery in the United States of
4	innovative cures, treatments, and preventive measures for
5	patients.
6	"SEC. 281B. DUTIES.
7	"For the purpose described in section 281A, the Coun-
8	cil shall—
9	"(1) foster collaboration and coordination among
10	the entities that comprise the Council, including aca-
11	demia, government agencies, industry, health care
12	payors and providers, patient advocates, and others
13	engaged in the cycle of discovery, development, and
14	delivery of life-saving and health-enhancing innova-
15	tive interventions;
16	"(2) undertake communication and dissemina-
17	tion activities;
18	"(3) publish information on the activities funded
19	under section 281D;
20	"(4) establish a strategic agenda for accelerating
21	the discovery, development, and delivery in the
22	United States of innovative cures, treatments, and
23	preventive measures for patients;
24	"(5) identify gaps and opportunities within and
25	across the discovery, development, and delivery cycle;

1	"(6) develop and propose recommendations based
2	on the gaps and opportunities so identified;
3	"(7) facilitate the interoperability of the compo-
4	nents of the discovery, development, and delivery
5	cycle;
6	"(8) propose recommendations that will facilitate
7	$precompetitive\ collaboration;$
8	"(9) identify opportunities to work with, but not
9	duplicate the efforts of, nonprofit organizations and
10	other public-private partnerships; and
11	"(10) identify opportunities for collaboration
12	with organizations operating outside of the United
13	States, such as the Innovative Medicines Initiative of
14	the European Union.
15	"SEC. 281C. ORGANIZATION; ADMINISTRATION.
16	"(a) Board of Directors.—
17	"(1) Establishment.—
18	"(A) In general.—The Council shall have
19	a Board of Directors (in this part referred to as
20	the 'Board of Directors'), which shall be com-
21	posed of the ex officio members under subpara-
22	graph (B) and the appointed members under
23	subparagraph (C). All members of the Board
24	shall be voting members.

1	"(B) Ex officio members.—The ex officio
2	members of the Board shall be the following indi-
3	viduals or their designees:
4	"(i) The Director of the National Insti-
5	tutes of Health.
6	"(ii) The Commissioner of Food and
7	Drugs.
8	"(iii) The Administrator of the Centers
9	for Medicare & Medicaid Services.
10	"(iv) The heads of five other Federal
11	agencies deemed by the Secretary to be en-
12	gaged in biomedical research and develop-
13	ment.
14	"(C) Appointed members.—The ap-
15	pointed members of the Board shall consist of 17
16	individuals, of whom—
17	"(i) 8 shall be appointed by the Comp-
18	troller General of the United States from a
19	list of nominations submitted by leading
20	trade associations—
21	"(I) 4 of whom shall be represent-
22	atives of the biopharmaceutical indus-
23	try;

1	"(II) 2 of whom shall be rep-
2	resentatives of the medical device in-
3	dustry; and
4	"(III) 2 of whom shall be rep-
5	resentatives of the information and
6	digital technology industry; and
7	"(ii) 9 shall be appointed by the
8	Comptroller General of the United States,
9	after soliciting nominations—
10	"(I) 2 of whom shall be represent-
11	atives of academic researchers;
12	"(II) 3 of whom shall be rep-
13	resentatives of patients;
14	"(III) 2 of whom shall be rep-
15	resentatives of health care providers;
16	and
17	"(IV) 2 of whom shall be rep-
18	resentatives of health care plans and
19	in surers.
20	"(D) Chair.—The Chair of the Board shall
21	be selected by the members of the Board by ma-
22	jority vote from among the members of the
23	Board.
24	"(2) Terms and vacancies.—

1	"(A) In General.—The term of office of
2	each member of the Board appointed under
3	paragraph (1)(C) shall be 5 years.
4	"(B) VACANCY.—Any vacancy in the mem-
5	bership of the Board—
6	"(i) shall not affect the power of the re-
7	maining members to execute the duties of
8	the Board; and
9	"(ii) shall be filled by appointment by
10	the appointed members described in para-
11	$graph\ (1)(C)\ by\ majority\ vote.$
12	"(C) Partial term.—If a member of the
13	Board does not serve the full term applicable
14	under subparagraph (A), the individual ap-
15	pointed under subparagraph (B) to fill the re-
16	sulting vacancy shall be appointed for the re-
17	mainder of the term of the predecessor of the in-
18	dividual.
19	"(3) Responsibilities.—Not later than 90 days
20	after the date on which the Council is incorporated
21	and its Board of Directors is fully constituted, the
22	Board of Directors shall establish bylaws and policies
23	for the Council that—
24	"(A) are published in the Federal Register
25	and available for public comment;

1	"(B) establish policies for the selection and,
2	as applicable, appointment of—
3	"(i) the officers, employees, agents, and
4	contractors of the Council; and
5	"(ii) the members of any committees of
6	the Council;
7	"(C) establish policies, including ethical
8	standards, for the conduct of programs and other
9	activities under section 281D; and
10	"(D) establish specific duties of the Execu-
11	tive Director.
12	"(4) Meetings.—
13	"(A) In General.—The Board of Directors
14	shall—
15	"(i) meet on a quarterly basis; and
16	"(ii) submit to Congress, and make
17	publicly available, the minutes of such meet-
18	ings.
19	"(B) AGENDA.—The Board of Directors
20	shall, not later than 3 months after the incorpo-
21	ration of the Council—
22	"(i) issue an agenda (in this part re-
23	ferred to as the 'agenda') outlining how the
24	Council will achieve the purpose described
25	in section 281A; and

1	"(ii) annually thereafter, in consulta-
2	tion with the Executive Director, review
3	and update such agenda.
4	"(b) Appointment and Incorporation.—Not later
5	than 6 months after the date of enactment of the 21st Cen-
6	tury Cures Act—
7	"(1) the Comptroller General of the United
8	States shall appoint the appointed members of the
9	Board of Directors under subsection $(a)(1)(C)$; and
10	"(2) the ex officio members of the Board of Di-
11	rectors under subsection $(a)(1)(B)$ shall serve as
12	incorporators and shall take whatever actions are nec-
13	essary to incorporate the Council.
14	"(c) Nonprofit Status.—In carrying out this part,
15	the Board of Directors shall establish such policies and by-
16	laws, and the Executive Director shall carry out such ac-
17	tivities, as may be necessary to ensure that the Council
18	maintains status as an organization that—
19	"(1) is described in subsection $(c)(3)$ of section
20	501 of the Internal Revenue Code of 1986; and
21	"(2) is, under subsection (a) of such section, ex-
22	empt from taxation.
23	"(d) Executive Director.—The Executive Director
24	shall—

"(1) be the chief executive officer of the Council; 1 2 and 3 "(2) subject to the oversight of the Board of Di-4 rectors, be responsible for the day-to-day management 5 of the Council. 6 "SEC. 281D. OPERATIONAL ACTIVITIES AND ASSISTANCE. 7 "(a) In General.—The Council shall establish a suffi-8 cient operational infrastructure to fulfill the duties specified in section 281B. 10 "(b) Private Sector Matching Funds.—The Council may accept financial or in-kind support from participating entities or private foundations or organizations when such support is deemed appropriate. 14 "SEC. 281E. TERMINATION; REPORT. 15 "(a) In General.—The Council shall terminate on September 30, 2023. 16 17 "(b) Report.—Not later than one year after the date on which the Council is established and each year thereafter, 18 19 the Executive Director shall submit to the appropriate congressional committees a report on the performance of the 21 Council. In preparing such report, the Council shall consult with a nongovernmental consultant with appropriate exper-

23 *tise*.

1	"SEC. 281F. FUNDING.
2	"For the each of fiscal years 2016 through 2023, there
3	is authorized to be appropriated \$10,000,000 to the Counci
4	for purposes of carrying out the duties of the Council under
5	this part.".
6	TITLE II—DEVELOPMENT
7	Subtitle A—Patient-Focused Drug
8	Development
9	SEC. 2001. DEVELOPMENT AND USE OF PATIENT EXPERI
10	ENCE DATA TO ENHANCE STRUCTURED RISK
11	BENEFIT ASSESSMENT FRAMEWORK.
12	(a) In General.—Section 505 of the Federal Food
13	Drug, and Cosmetic Act (21 U.S.C. 355) is amended—
14	(1) in subsection (d), by striking "The Secretary
15	shall implement" and all that follows through "pre
16	market approval of a drug."; and
17	(2) by adding at the end the following new sub-
18	sections:
19	"(x) Structured Risk-Benefit Assessment
20	Framework.—
21	"(1) In General.—The Secretary shall imple
22	ment a structured risk-benefit assessment framework
23	in the new drug approval process—
24	"(A) to facilitate the balanced consideration
25	of henefits and risks: and

1	"(B) to develop and implement a consistent
2	and systematic approach to the discussion of,
3	regulatory decisionmaking with respect to, and
4	the communication of, the benefits and risks of
5	new drugs.
6	"(2) Rule of construction.—Nothing in
7	paragraph (1) shall alter the criteria for evaluating
8	an application for premarket approval of a drug.
9	"(y) Development and Use of Patient Experi-
10	ENCE DATA TO ENHANCE STRUCTURED RISK-BENEFIT AS-
11	SESSMENT FRAMEWORK.—
12	"(1) In general.—Not later than two years
13	after the date of the enactment of this subsection, the
14	Secretary shall establish and implement processes
15	under which—
16	"(A) an entity seeking to develop patient ex-
17	perience data may submit to the Secretary—
18	"(i) initial research concepts for feed-
19	back from the Secretary; and
20	"(ii) with respect to patient experience
21	data collected by the entity, draft guidance
22	documents, completed data, and summaries
23	and analyses of such data;

1	"(B) the Secretary may request such an en-
2	tity to submit such documents, data, and sum-
3	maries and analyses; and
4	"(C) patient experience data may be devel-
5	oped and used to enhance the structured risk-
6	benefit assessment framework under subsection
7	(x).
8	"(2) Patient experience data.—In this sub-
9	section, the term 'patient experience data' means data
10	collected by patients, parents, caregivers, patient ad-
11	vocacy organizations, disease research foundations,
12	medical researchers, research sponsors, or other par-
13	ties determined appropriate by the Secretary that is
14	intended to facilitate or enhance the Secretary's risk-
15	benefit assessments, including information about the
16	impact of a disease or a therapy on patients' lives.".
17	(b) Guidance.—
18	(1) In General.—The Secretary of Health and
19	Human Services shall publish guidance on the imple-
20	mentation of subsection (y) of section 505 of the Fed-
21	eral Food, Drug, and Cosmetic Act (21 U.S.C. 355),
22	as added by subsection (a). Such guidance shall in-
23	clude—
24	(A) with respect to draft guidance docu-
25	ments, data, or summaries and analyses sub-

1	mitted to the Secretary under paragraph $(1)(A)$
2	of such subsection, guidance—
3	(i) specifying the timelines for the re-
4	view of such documents, data, or summaries
5	and analyses by the Secretary; and
6	(ii) on how the Secretary will use such
7	documents, data, or summaries and anal-
8	yses to update any guidance documents
9	published under this subsection or publish
10	new guidance;
11	(B) with respect to the collection and anal-
12	ysis of patient experience data (as defined in
13	paragraph (2) of such subsection (y)), guidance
14	on—
15	(i) methodological considerations for
16	the collection of patient experience data,
17	which may include structured approaches to
18	gathering information on—
19	(I) the experience of a patient liv-
20	ing with a particular disease;
21	(II) the burden of living with or
22	managing the disease;
23	(III) the impact of the disease on
24	daily life and long-term functioning;
25	and

1	(IV) the effect of current thera-
2	peutic options on different aspects of
3	the disease; and
4	(ii) the establishment and maintenance
5	of registries designed to increase under-
6	standing of the natural history of a disease;
7	(C) methodological approaches that may be
8	used to assess patients' beliefs with respect to the
9	benefits and risks in the management of the pa-
10	tient's disease; and
11	(D) methodologies, standards, and potential
12	experimental designs for patient-reported out-
13	comes.
14	(2) Timing.—Not later than 3 years after the
15	date of the enactment of this Act, the Secretary of
16	Health and Human Services shall issue draft guid-
17	ance on the implementation of subsection (y) of sec-
18	tion 505 of the Federal Food, Drug, and Cosmetic Act
19	(21 U.S.C. 355), as added by subsection (a). The Sec-
20	retary shall issue final guidance on the implementa-
21	tion of such subsection not later than one year after
22	the date on which the comment period for the draft
23	guidance closes.
24	(3) Workshops.—

1	(A) In general.—Not later than 6 months
2	after the date of the enactment of this Act and
3	once every 6 months during the following 12-
4	month period, the Secretary of Health and
5	Human Services shall convene a workshop to ob-
6	tain input regarding methodologies for devel-
7	oping the guidance under paragraph (1), includ-
8	ing the collection of patient experience data.
9	(B) Attendees.—A workshop convened
10	under this paragraph shall include—
11	(i) patients;
12	(ii) representatives from patient advo-
13	cacy organizations, biopharmaceutical com-
14	panies, and disease research foundations;
15	(iii) representatives of the reviewing
16	divisions of the Food and Drug Administra-
17	$tion; \ and$
18	(iv) methodological experts with sig-
19	nificant expertise in patient experience
20	data.
21	(4) Public meeting.—Not later than 90 days
22	after the date on which the draft guidance is pub-
23	lished under this subsection, the Secretary of Health
24	and Human Services shall convene a public meeting
25	to solicit input on the guidance.

Subtitle B—Qualification and Use 1 of Drug Development Tools 2 SEC. 2021. QUALIFICATION OF DRUG DEVELOPMENT TOOLS. (a) FINDINGS.—Congress finds the following: 4 5 (1) Development of new drugs has become in-6 creasingly challenging and resource intensive. 7 (2) Development of drug development tools can 8 benefit the availability of new medical therapies by 9 helping to translate scientific discoveries into clinical 10 applications. 11 (3) Biomedical research consortia (as defined in 12 section 507(f) of the Federal Food, Drug, and Cos-13 metic Act, as added by subsection (c)) can play a val-14 uable role in helping to develop and qualify drug de-15 velopment tools. 16 (b) Sense of Congress.—It is the sense of Congress 17 that— 18 (1) Congress should promote and facilitate a collaborative effort among the biomedical research con-19 20 sortia described in subsection (a)(3)— 21 (A) to develop, through a transparent public 22 process, data standards and scientific approaches 23 to data collection accepted by the medical and

clinical research community for purposes of

qualifying drug development tools;

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1	(B) to coordinate efforts toward developing
2	and qualifying drug development tools in key
3	therapeutic areas; and
4	(C) to encourage the development of acces-
5	sible databases for collecting relevant drug devel-
6	opment tool data for such purposes; and
7	(2) an entity seeking to qualify a drug develop-
8	ment tool should be encouraged, in addition to con-
9	sultation with the Secretary, to consult with bio-
10	medical research consortia and other individuals and
11	entities with expert knowledge and insights that may
12	assist the requestor and benefit the process for such
13	qualification.
14	(c) Qualification of Drug Development Tools.—
15	Chapter V of the Federal Food, Drug, and Cosmetic Act
16	is amended by inserting after section 506F the following
17	new section:
18	"SEC. 507. QUALIFICATION OF DRUG DEVELOPMENT TOOLS.
19	"(a) Process for Qualification.—
20	"(1) In general.—The Secretary shall establish
21	a process for the qualification of drug development
22	tools for a proposed context of use under which—
23	"(A)(i) a requestor initiates such process by
24	submitting a letter of intent to the Secretary;
25	and

1	"(ii) the Secretary shall accept or decline to
2	accept such letter of intent;
3	"(B)(i) if the Secretary accepts the letter of
4	intent, a requestor shall submit a qualification
5	plan to the Secretary; and
6	"(ii) the Secretary shall accept or decline to
7	accept the qualification plan; and
8	" $(C)(i)$ if the Secretary accepts the quali-
9	fication plan, the requestor submits to the Sec-
10	retary a full qualification package;
11	"(ii) the Secretary shall determine whether
12	to accept such qualification package for review;
13	and
14	"(iii) if the Secretary accepts such quali-
15	fication package for review, the Secretary shall
16	conduct such review in accordance with this sec-
17	tion.
18	"(2) Acceptance and review of submis-
19	SIONS.—
20	"(A) In general.—The succeeding provi-
21	sions of this paragraph shall apply with respect
22	to the treatment of a letter of intent, a qualifica-
23	tion plan, or a full qualification package sub-
24	mitted under paragraph (1) (referred to in this
25	paragraph as 'qualification submissions').

1	"(B) Acceptance factors; nonaccept-
2	ANCE.—The Secretary shall determine whether to
3	accept a qualification submission based on fac-
4	tors which may include the scientific merit of the
5	submission and the available resources of the
6	Food and Drug Administration to review the
7	qualification submission. A determination not to
8	accept a submission under paragraph (1) shall
9	not be construed as a final determination by the
10	Secretary under this section regarding the quali-
11	fication of a drug development tool for its pro-
12	posed context of use.
13	"(C) Prioritization of qualification
14	REVIEW.—The Secretary may prioritize the re-
15	view of a full qualification package submitted
16	under paragraph (1) with respect to a drug de-
17	velopment tool, based on factors determined ap-
18	propriate by the Secretary, including—
19	"(i) as applicable, the severity, rarity,
20	or prevalence of the disease or condition
21	targeted by the drug development tool and
22	the availability or lack of alternative treat-
23	ments for such disease or condition; and
24	"(ii) the identification, by the Sec-
25	retary or by biomedical research consortia

1	and other expert stakeholders, of such a
2	drug development tool and its proposed con-
3	text of use as a public health priority.
4	"(D) Engagement of external ex-
5	PERTS.—The Secretary may, for purposes of the
6	review of qualification submissions, through the
7	use of cooperative agreements, grants, or other
8	appropriate mechanisms, consult with bio-
9	medical research consortia and may consider the
10	recommendations of such consortia with respect
11	to the review of any qualification plan submitted
12	under paragraph (1) or the review of any full
13	qualification package under paragraph (3).
14	"(3) Review of full qualification pack-
15	AGE.—The Secretary shall—
16	"(A) conduct a comprehensive review of a
17	full qualification package accepted under para-
18	$graph\ (1)(C);\ and$
19	"(B) determine whether the drug develop-
20	ment tool at issue is qualified for its proposed
21	context of use.
22	"(4) Qualification.—The Secretary shall deter-
23	mine whether a drug development tool is qualified for
24	a proposed context of use based on the scientific merit

1	of a full qualification package reviewed under para-
2	graph(3).
3	"(b) Effect of Qualification.—
4	"(1) In general.—A drug development tool de-
5	termined to be qualified under subsection (a)(4) for a
6	proposed context of use specified by the requestor may
7	be used by any person in such context of use for the
8	purposes described in paragraph (2).
9	"(2) Use of a drug development tool.—
10	Subject to paragraph (3), a drug development tool
11	qualified under this section may be used for—
12	"(A) supporting or obtaining approval or
13	licensure (as applicable) of a drug or biological
14	product (including in accordance with section
15	506(c)) under section 505 of this Act or section
16	351 of the Public Health Service Act; or
17	"(B) supporting the investigational use of a
18	drug or biological product under section 505(i)
19	of this Act or section 351(a)(3) of the Public
20	Health Service Act.
21	"(3) Rescission or modification.—
22	"(A) In general.—The Secretary may re-
23	scind or modify a determination under this sec-
24	tion to qualify a drug development tool if the
25	Secretary determines that the drug development

1	tool is not appropriate for the proposed context
2	of use specified by the requestor. Such a deter-
3	mination may be based on new information that
4	calls into question the basis for such qualifica-
5	tion.
6	"(B) Meeting for review.—If the Sec-
7	retary rescinds or modifies under subparagraph
8	(A) a determination to qualify a drug develop-
9	ment tool, the requestor involved shall be granted
10	a request for a meeting with the Secretary to dis-
11	cuss the basis of the Secretary's decision to re-
12	scind or modify the determination before the ef-
13	fective date of the rescission or modification.
14	"(c) Transparency.—
15	"(1) In general.—Subject to paragraph (3), the
16	Secretary shall make publicly available, and update
17	on at least a biannual basis, on the Internet website
18	of the Food and Drug Administration the following:
19	"(A) Information with respect to each qual-
20	ification submission under the qualification
21	process under subsection (a), including—
22	"(i) the stage of the review process ap-
23	plicable to the submission;
24	"(ii) the date of the most recent change
25	$in\ stage\ status;$

1	"(iii) whether the external scientific ex-
2	perts were utilized in the development of a
3	qualification plan or the review of a full
4	qualification package; and
5	"(iv) submissions from requestors
6	under the qualification process under sub-
7	section (a), including any data and evi-
8	dence contained in such submissions, and
9	any updates to such submissions.
10	"(B) The Secretary's formal written deter-
11	minations in response to such qualification sub-
12	missions.
13	"(C) Any rescissions or modifications under
14	subsection (b)(3) of a determination to qualify a
15	drug development tool.
16	"(D) Summary reviews that document con-
17	clusions and recommendations for determina-
18	tions to qualify drug development tools under
19	subsection (a).
20	"(E) A comprehensive list of—
21	"(i) all drug development tools quali-
22	fied under subsection (a); and
23	"(ii) all surrogate endpoints which
24	were the basis of approval or licensure (as
25	applicable) of a drug or biological product

1	(including in accordance with section
2	506(c)) under section 505 of this Act or sec-
3	tion 351 of the Public Health Service Act.
4	"(2) Relation to trade secrets act.—Infor-
5	mation made publicly available by the Secretary
6	under paragraph (1) shall be considered a disclosure
7	authorized by law for purposes of section 1905 of title
8	18, United States Code.
9	"(3) Applicability.—Nothing in this section
10	shall be construed as authorizing the Secretary to dis-
11	close any information contained in an application
12	submitted under section 505 of this Act or section 351
13	of the Public Health Service Act that is confidential
14	commercial or trade secret information subject to sec-
15	tion 552(b)(4) of title 5, United States Code, or sec-
16	tion 1905 of title 18, United States Code.
17	"(d) Rule of Construction.—Nothing in this sec-
18	tion shall be construed—
19	"(1) to alter the standards of evidence under sub-
20	section (c) or (d) of section 505, including the sub-
21	stantial evidence standard in such subsection (d), or
22	under section 351 of the Public Health Service Act (as
23	applicable); or
24	"(2) to limit the authority of the Secretary to
25	approve or license products under this Act or the Pub-

- 1 lic Health Service Act, as applicable (as in effect be-
- 2 fore the date of the enactment of the 21st Century
- 3 Cures Act).
- 4 "(e) AUTHORIZATION OF APPROPRIATIONS.—There are
- 5 authorized to be appropriated to carry out this section,
- 6 \$10,000,000 for each of fiscal years 2016 through 2020.
- 7 "(f) Definitions.—In this section:
- 8 "(1) BIOMARKER.—(A) The term 'biomarker'
 9 means a characteristic (such as a physiologic,
 10 pathologic, or anatomic characteristic or measure11 ment) that is objectively measured and evaluated as
 12 an indicator of normal biologic processes, pathologic
 13 processes, or biological responses to a therapeutic
 14 intervention: and
 - "(B) such term includes a surrogate endpoint.
- "(2) BIOMEDICAL RESEARCH CONSORTIA.—The 16 17 term 'biomedical research consortia' means collabo-18 rative groups that may take the form of public-pri-19 vate partnerships and may include government agen-20 cies, institutions of higher education (as defined in 21 section 101(a) of the Higher Education Act of 1965, 22 patient advocacy groups, industry representatives, 23 clinical and scientific experts, and other relevant entities and individuals. 24

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1	"(3) Clinical outcome assessment.—(A) The
2	term 'clinical outcome assessment' means a measure-
3	ment of a patient's symptoms, overall mental state, or
4	the effects of a disease or condition on how the patient
5	functions; and
6	"(B) such term includes a patient-reported out-
7	come.
8	"(4) Context of use.—The term 'context of
9	use' means, with respect to a drug development tool,
10	a statement that describes the circumstances under
11	which the drug development tool is to be used in drug
12	development and regulatory review.
13	"(5) Drug development tool.—The term
14	'drug development tool' includes—
15	"(A) a biomarker;
16	"(B) a clinical outcome assessment; and
17	"(C) any other method, material, or meas-
18	ure that the Secretary determines aids drug de-
19	velopment and regulatory review for purposes of
20	$this\ section.$
21	"(6) Patient-reported outcome.—The term
22	'patient-reported outcome' means a measurement
23	based on a report from a patient regarding the status
24	of the patient's health condition without amendment

- or interpretation of the patient's report by a clinician or any other person.
 - "(7) QUALIFICATION.—The terms 'qualification' and 'qualified' mean a determination by the Secretary that a drug development tool and its proposed context of use can be relied upon to have a specific interpretation and application in drug development and regulatory review under this Act.
 - "(8) REQUESTOR.—The term 'requestor' means an entity or entities, including a drug sponsor or a biomedical research consortia, seeking to qualify a drug development tool for a proposed context of use under this section.
 - "(9) Surrogate endpoint end as a marker, such as a laboratory measurement, radiographic image, physical sign, or other measure, that is not itself a direct measurement of clinical benefit, and—
 - "(A) is known to predict clinical benefit and could be used to support traditional approval of a drug or biological product; or
 - "(B) is reasonably likely to predict clinical benefit and could be used to support the accelerated approval of a drug or biological product in accordance with section 506(c).".

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1	(d) Guidance.—
2	(1) In General.—The Secretary of Health and
3	Human Services shall, in consultation with bio-
4	medical research consortia (as defined in subsection
5	(f) of section 507 the Federal Food, Drug, and Cos-
6	metic Act (as added by subsection (c))) and other in-
7	terested parties through a collaborative public process,
8	issue guidance to implement such section 507 that—
9	(A) provides a conceptual framework de-
10	scribing appropriate standards and scientific
11	approaches to support the development of bio-
12	markers delineated under the taxonomy estab-
13	lished under paragraph (3);
14	(B) makes recommendations for dem-
15	onstrating that a surrogate endpoint is reason-
16	ably likely to predict clinical benefit for the pur-
17	pose of supporting the accelerated approval of a
18	drug under section 506(c) of the Federal Food,
19	Drug, and Cosmetic Act (21 U.S.C. 356(c));
20	(C) with respect to the qualification process
21	under such section 507—
22	(i) describes the requirements that enti-
23	ties seeking to qualify a drug development
24	tool under such section shall observe when
25	engaging in such process;

1	(ii) outlines reasonable timeframes for
2	the Secretary's review of letters, qualifica-
3	tion plans, or full qualification packages
4	submitted under such process; and
5	(iii) establishes a process by which
6	such entities or the Secretary may consult
7	with biomedical research consortia and
8	other individuals and entities with expert
9	knowledge and insights that may assist the
10	Secretary in the review of qualification
11	plans and full qualification submissions
12	under such section; and
13	(D) includes such other information as the
14	Secretary determines appropriate.
15	(2) Timing.—Not later than 24 months after the
16	date of the enactment of this Act, the Secretary of
17	Health and Human Services shall issue draft guid-
18	ance under paragraph (1) on the implementation of
19	section 507 of the Federal Food, Drug, and Cosmetic
20	Act (as added by subsection (c)). The Secretary shall
21	issue final guidance on the implementation of such
22	section not later than 6 months after the date on
23	which the comment period for the draft guidance
24	closes.
25	(3) Taxonomy.—

- (A) In General.—For purposes of informing guidance under this subsection, the Secretary of Health and Human Services shall, in consultation with biomedical research consortia and other interested parties through a collaborative public process, establish a taxonomy for the classification of biomarkers (and related scientific concepts) for use in drug development.
 - (B) Public Availability.—Not later than 12 months after the date of the enactment of this Act, the Secretary of Health and Human Services shall make such taxonomy publicly available in draft form for public comment. The Secretary shall finalize the taxonomy not later than 12 months after the close of the public comment period.

(e) Meeting and Report.—

(1) MEETING.—Not later than 12 months after the date of the enactment of this Act, the Secretary of Health and Human Services shall convene a public meeting to describe and solicit public input regarding the qualification process under section 507 of the Federal Food, Drug, and Cosmetic Act, as added by subsection (c).

1	(2) Report.—Not later than 5 years after the
2	date of the enactment of this Act, the Secretary shall
3	make publicly available on the Internet website of the
4	Food and Drug Administration a report. Such report
5	shall include, with respect to the qualification process
6	under section 507 of the Federal Food, Drug, and
7	Cosmetic Act, as added by subsection (c), information
8	on—
9	(A) the number of requests submitted, as a
10	letter of intent, for qualification of a drug devel-
11	opment tool (as defined in subsection (f) of such
12	section);
13	(B) the number of such requests accepted
14	and determined to be eligible for submission of a
15	qualification plan or full qualification package
16	(as such terms are defined in such subsection),
17	respectively;
18	(C) the number of such requests for which
19	external scientific experts were utilized in the de-
20	velopment of a qualification plan or review of a
21	full qualification package; and
22	(D) the number of qualification plans and
23	full qualification packages, respectively, sub-
24	mitted to the Secretary; and

1	(3) the drug development tools qualified through
2	such qualification process, specified by type of tool,
3	such as a biomarker or clinical outcome assessment
4	(as such terms are defined in subsection (f) of such
5	section 507).
6	SEC. 2022. ACCELERATED APPROVAL DEVELOPMENT PLAN
7	(a) In General.—Section 506 of the Federal Food,
8	Drug, and Cosmetic Act (21 U.S.C. 356) is amended by
9	adding the following subsection:
10	"(g) Accelerated Approval Development
11	PLAN.—
12	"(1) In general.—In the case of a drug that
13	the Secretary determines may be eligible for acceler-
14	ated approval in accordance with subsection (c), the
15	sponsor of such drug may request, at any time after
16	the submission of an application for the investigation
17	of the drug under section 505(i) of this Act or section
18	351(a)(3) of the Public Health Service Act, that the
19	Secretary agree to an accelerated approval develop-
20	ment plan described in paragraph (2).
21	"(2) Plan described in
22	this paragraph, with respect to a drug described in
23	paragraph (1), is an accelerated approval develop-
24	ment plan, which shall include agreement on—

1	"(A) the surrogate endpoint to be assessed
2	under such plan;
3	"(B) the design of the study that will utilize
4	the surrogate endpoint; and
5	"(C) the magnitude of the effect of the drug
6	on the surrogate endpoint that is the subject of
7	the agreement that would be sufficient to form
8	the primary basis of a claim that the drug is ef-
9	fective.
10	"(3) Modification; termination.—The Sec-
11	retary may require the sponsor of a drug that is the
12	subject of an accelerated approval development plan
13	to modify or terminate the plan if additional data or
14	information indicates that—
15	"(A) the plan as originally agreed upon is
16	no longer sufficient to demonstrate the safety and
17	effectiveness of the drug involved; or
18	"(B) the drug is no longer eligible for accel-
19	erated approval under subsection (c).
20	"(4) Sponsor consultation.—If the Secretary
21	requires the modification or termination of an accel-
22	erated approval development plan under paragraph
23	(3), the sponsor shall be granted a request for a meet-
24	ing to discuss the basis of the Secretary's decision be-

1	fore the effective date of the modification or termi-
2	nation.
3	"(5) Definition.—In this section, the term 'ac-
4	celerated approval development plan' means a devel-
5	opment plan agreed upon by the Secretary and the
6	sponsor submitting the plan that contains study pa-
7	rameters for the use of a surrogate endpoint that—
8	"(A) is reasonably likely to predict clinical
9	benefit; and
10	"(B) is intended to be the basis of the accel-
11	erated approval of a drug in accordance with
12	subsection (c).".
13	(b) Technical Amendments.—Section 506 of the
14	Federal Food, Drug, and Cosmetic Act (21 U.S.C. 356) is
15	amended—
16	(1) by striking "(f) Awareness Efforts" and
17	inserting "(e) AWARENESS EFFORTS"; and
18	(2) by striking "(e) Construction" and insert-
19	ing "(f) Construction".

1	Subtitle C—FDA Advancement of
2	Precision Medicine
3	SEC. 2041. PRECISION MEDICINE GUIDANCE AND OTHER
4	PROGRAMS OF FOOD AND DRUG ADMINISTRA-
5	TION.
6	Chapter V of the Federal Food, Drug, and Cosmetic
7	Act (21 U.S.C. 351 et seq.) is amended by adding at the
8	end the following:
9	$"Subchapter J-Precision \ Medicine$
10	"SEC. 591. GENERAL AGENCY GUIDANCE ON PRECISION
11	MEDICINE.
12	"(a) In General.—The Secretary shall issue and pe-
13	riodically update guidance to assist sponsors in the develop-
14	ment of a precision drug or biological product. Such guid-
15	ance shall—
16	"(1) define the term 'precision drug or biological
17	product'; and
18	"(2) address the topics described in subsection
19	<i>(b)</i> .
20	"(b) Certain Issues.—The topics to be addressed by
21	guidance under subsection (a) are—
22	"(1) the evidence needed to support the use of
23	biomarkers (as defined in section 507(e)) that identify
24	subsets of patients as likely responders to therapies in
25	order to streamline the conduct of clinical trials:

1	"(2) recommendations for the design of studies to
2	demonstrate the validity of a biomarker as a pre-
3	dictor of drug or biological product response;
4	"(3) the manner and extent to which a benefit-
5	risk assessment may be affected when clinical trials
6	are limited to patient population subsets that are
7	identified using biomarkers;
8	"(4) the development of companion diagnostics
9	in the context of a drug development program; and
10	"(5) considerations for developing biomarkers
11	that inform prescribing decisions for a drug or bio-
12	logical product, and when information regarding a
13	biomarker may be included in the approved prescrip-
14	tion labeling for a precision drug or biological prod-
15	uct.
16	"(c) Date Certain for Initial Guidance.—The
17	Secretary shall issue guidance under subsection (a) not
18	later than 18 months after the date of the enactment of the
19	21st Century Cures Act.
20	"SEC. 592. PRECISION MEDICINE REGARDING ORPHAN-
21	DRUG AND EXPEDITED-APPROVAL PRO-
22	GRAMS.
23	"(a) In General.—In the case of a precision drug
24	or biological product that is the subject of an application
25	submitted under section 505(b)(1), or section 351(a) of the

1	Public Health Service Act, for the treatment of a serious
2	or life-threatening disease or condition and has been des-
3	ignated under section 526 as a drug for a rare disease or
4	condition, the Secretary may—
5	"(1) consistent with applicable standards for ap-
6	proval, rely upon data or information previously sub-
7	mitted by the sponsor of the precision drug or biologi-
8	cal product, or another sponsor, provided that the
9	sponsor of the precision drug or biological product
10	has obtained a contractual right of reference to such
11	other sponsor's data and information, in an applica-
12	tion approved under section 505(c) or licensed under
13	section 351(a) of the Public Health Service Act, as
14	applicable—
15	"(A) for a different drug or biological prod-
16	uct; or
17	"(B) for a different indication for such pre-
18	cision drug or biological product,
19	in order to expedite clinical development for a preci-
20	sion drug or biological product that is using the same
21	or similar approach as that used to support approval
22	of the prior approved application or license, as ap-
23	propriate; and
24	"(2) as appropriate, consider the application for
25	approval of such precision drug or biological product

1	to be eligible for expedited review and approval pro-
2	grams described in section 506, including accelerated
3	approval in accordance with subsection (c) of such
4	section.
5	"(b) Rule of Construction.—Nothing in this sec-
6	tion shall be construed to—
7	"(1) limit the authority of the Secretary to ap-
8	prove products pursuant to this Act and the Public
9	Health Service Act as authorized prior to the date of
10	enactment of this section; or
11	"(2) confer any new rights, beyond those author-
12	ized under this Act prior to enactment of this section,
13	with respect to a sponsor's ability to reference infor-
14	mation contained in another application submitted
15	under section 505(b)(1) of this Act or section 351(a)
16	of the Public Health Service Act.".
17	Subtitle D—Modern Trial Design
18	and Evidence Development
19	SEC. 2061. BROADER APPLICATION OF BAYESIAN STATIS-
20	TICS AND ADAPTIVE TRIAL DESIGNS.
21	(a) Proposals for Use of Innovative Statistical
22	Methods in Clinical Protocols for Drugs and Bio-
23	LOGICAL PRODUCTS.—For purposes of assisting sponsors in
24	incorporating adaptive trial design and Bayesian methods
25	into proposed clinical protocols and applications for new

1	drugs under section 505 of the Federal Food, Drug, and
2	Cosmetic Act (21 U.S.C. 355) and biological products under
3	section 351 of the Public Health Service Act (42 U.S.C.
4	262), the Secretary shall conduct a public meeting and issue
5	guidance in accordance with subsection (b).
6	(b) Guidance Addressing Use of Adaptive Trial
7	Designs and Bayesian Methods.—
8	(1) In General.—The Secretary of Health and
9	Human Services, acting through the Commissioner of
10	Food and Drugs (in this subsection referred to as the
11	"Secretary"), shall—
12	(A) update and finalize the draft guidance
13	addressing the use of adaptive trial design for
14	drugs and biological products; and
15	(B) issue draft guidance on the use of
16	Bayesian methods in the development and regu-
17	latory review and approval or licensure of drugs
18	and biological products.
19	(2) Contents.—The guidances under paragraph
20	(1) shall address—
21	(A) the use of adaptive trial designs and
22	Bayesian methods in clinical trials, including
23	clinical trials proposed or submitted to help to
24	satisfy the substantial evidence standard under

1	section $505(d)$ of the Federal Food, Drug, and
2	Cosmetic Act (21 U.S.C. $355(d)$);
3	(B) how sponsors may obtain feedback from
4	the Secretary on technical issues related to mod-
5	eling and simulations prior to—
6	(i) completion of such modeling or sim-
7	ulations; or
8	(ii) the submission of resulting infor-
9	mation to the Secretary;
10	(C) the types of quantitative and qualitative
11	information that should be submitted for review;
12	and
13	(D) recommended analysis methodologies.
14	(3) Public meeting.—Prior to updating or de-
15	veloping the guidances required by paragraph (1), the
16	Secretary shall consult with stakeholders, including
17	representatives of regulated industry, academia, pa-
18	tient advocacy organizations, and disease research
19	foundations, through a public meeting to be held not
20	later than 1 year after the date of enactment of this
21	Act.
22	(4) Schedule.—The Secretary shall publish—
23	(A) the final guidance required by para-
24	graph (1)(A) not later than 18 months after the

1	date of the public meeting required by paragraph
2	(3); and
3	(B) the guidance required by paragraph
4	(1)(B) not later than 48 months after the date of
5	the public meeting required by paragraph (3).
6	SEC. 2062. UTILIZING EVIDENCE FROM CLINICAL EXPERI-
7	ENCE.
8	Chapter V of the Federal Food, Drug, and Cosmetic
9	Act is amended by inserting after section 505E of such Act
10	(21 U.S.C. 355f) the following:
11	"SEC. 505F. UTILIZING EVIDENCE FROM CLINICAL EXPERI-
12	ENCE.
13	"(a) In General.—The Secretary shall establish a
14	program to evaluate the potential use of evidence from clin-
15	ical experience—
16	"(1) to help to support the approval of a new in-
17	dication for a drug approved under section 505(b);
18	and
19	"(2) to help to support or satisfy postapproval
20	study requirements.
21	"(b) Evidence From Clinical Experience De-
22	FINED.—In this section, the term 'evidence from clinical ex-
23	perience' means data regarding the usage, or the potential
24	benefits or risks, of a drug derived from sources other than

1	randomized clinical trials, including from observational
2	studies, registries, and therapeutic use.
3	"(c) Program Framework.—
4	"(1) In general.—Not later than 18 months
5	after the date of enactment of this section, the Sec-
6	retary shall establish a draft framework for imple-
7	mentation of the program under this section.
8	"(2) Contents of Framework.—The frame-
9	work shall include information describing—
10	"(A) the current sources of data developed
11	through clinical experience, including ongoing
12	safety surveillance, registry, claims, and patient-
13	centered outcomes research activities;
14	"(B) the gaps in current data collection ac-
15	tivities;
16	"(C) the current standards and methodolo-
17	gies for collection and analysis of data generated
18	through clinical experience; and
19	"(D) the priority areas, remaining chal-
20	lenges, and potential pilot opportunities that the
21	program established under this section will ad-
22	dress.
23	"(3) Consultation.—
24	"(A) In General.—In developing the pro-
25	gram framework under this subsection, the Sec-

1	retary shall consult with regulated industry, aca-
2	demia, medical professional organizations, rep-
3	resentatives of patient advocacy organizations,
4	disease research foundations, and other interested
5	parties.
6	"(B) Process.—The consultation under
7	subparagraph (A) may be carried out through
8	approaches such as—
9	"(i) a public-private partnership with
10	the entities described in such subparagraph
11	in which the Secretary may participate; or
12	"(ii) a contract, grant, or other ar-
13	rangement, as determined appropriate by
14	the Secretary with such a partnership or an
15	independent research organization.
16	"(d) Program Implementation.—The Secretary
17	shall, not later than 24 months after the date of enactment
18	of this section and in accordance with the framework estab-
19	lished under subsection (c), implement the program to
20	evaluate the potential use of evidence from clinical experi-
21	ence.
22	"(e) Guidance for Industry.—The Secretary
23	shall—
24	"(1) utilize the program established under sub-
25	section (a), its activities, and any subsequent pilots or

1	written reports, to inform a guidance for industry
2	on—
3	"(A) the circumstances under which spon-
4	sors of drugs and the Secretary may rely on evi-
5	dence from clinical experience for the purposes
6	described in subsection (a)(1) or (a)(2); and
7	"(B) the appropriate standards and meth-
8	odologies for collection and analysis of evidence
9	from clinical experience submitted for such pur-
10	poses;
11	"(2) not later than 36 months after the date of
12	enactment of this section, issue draft guidance for in-
13	dustry as described in paragraph (1); and
14	"(3) not later than 48 months after the date of
15	enactment of this section, after providing an oppor-
16	tunity for public comment on the draft guidance,
17	issue final guidance.
18	"(f) Rule of Construction.—
19	"(1) Subject to paragraph (2), nothing in this
20	section prohibits the Secretary from using evidence
21	from clinical experience for purposes not specified in
22	this section, provided the Secretary determines that
23	sufficient basis exists for any such nonspecified use.
24	"(2) This section shall not be construed to
25	alter—

1	"(A) the standards of evidence under—
2	"(i) subsection (c) or (d) of section 505,
3	including the substantial evidence standard
4	in such subsection (d); or
5	"(ii) section 351(a) of the Public
6	Health Service Act; or
7	"(B) the Secretary's authority to require
8	postapproval studies or clinical trials, or the
9	standards of evidence under which studies or
10	trials are evaluated.
11	"SEC. 505G. COLLECTING EVIDENCE FROM CLINICAL EXPE-
12	RIENCE THROUGH TARGETED EXTENSIONS
13	OF THE SENTINEL SYSTEM.
14	"(a) In General.—The Secretary shall, in parallel to
15	$implementing\ the\ program\ established\ under\ section\ 505F$
16	and in order to build capacity for utilizing the evidence
17	from clinical experience described in that section, identify
18	and execute pilot demonstrations to extend existing use of
19	$the \ Sentinel \ System \ surveillance \ infrastructure \ authorized$
20	$under\ section\ 505(k).$
21	"(b) Pilot Demonstrations.—
22	"(1) In General.—The Secretary—
23	"(A) shall design and implement pilot dem-
24	onstrations to utilize data captured through the
25	Sentinel System surveillance infrastructure au-

1	thorized under section 505(k) for purposes of, as
2	appropriate—
3	"(i) generating evidence from clinical
4	experience to improve characterization or
5	assessment of risks or benefits of a drug ap-
6	$proved\ under\ section\ 505(c);$
7	"(ii) protecting the public health; or
8	"(iii) advancing patient-centered care;
9	and
10	"(B) may make strategic linkages with
11	sources of complementary public health data and
12	infrastructure the Secretary determines appro-
13	priate and necessary.
14	"(2) Consultation.—In developing the pilot
15	demonstrations under this subsection, the Secretary
16	shall—
17	"(A) consult with regulated industry, aca-
18	demia, medical professional organizations, rep-
19	resentatives of patient advocacy organizations,
20	disease research foundations, and other interested
21	parties through a public process; and
22	"(B) develop a framework to promote ap-
23	propriate transparency and dialogue about re-
24	search conducted under these pilot demonstra-
25	tions, including by—

1	"(i) providing adequate notice to a
2	sponsor of a drug approved under section
3	505 or section 351 of the Public Health
4	Service Act of the Secretary's intent to con-
5	duct analyses of such sponsor's drug or
6	drugs under these pilot demonstrations;
7	"(ii) providing adequate notice of the
8	findings related to analyses described in
9	clause (i) and an opportunity for the spon-
10	sor of such drug or drugs to comment on
11	such findings; and
12	"(iii) ensuring the protection from
13	public disclosure of any information that is
14	a trade secret or confidential information
15	subject to section 552(b)(4) of title 5, United
16	States Code, or section 1905 of title 18,
17	United States Code.
18	"(3) Public Health Exemption.—The Sec-
19	retary may—
20	"(A) deem such pilot demonstrations public
21	health activities, permitting the use and disclo-
22	sure of protected health information as described
23	in section 164.512(b)(1)(iii) of title 45, Code of
24	Federal Regulations (or any successor regula-
25	tion) and exempted as a public health activity as

1 described in section 46.101(b)(5) of title 46, Code 2 of Federal Regulations (or any successor regula-3 tion); and

"(B) deem safety surveillance performed at the request of the Food and Drug Administration or under such jurisdiction by a sponsor with responsibility for a drug approved under this section or section 351 of the Public Health Services Act using the Sentinel System surveillance infrastructure authorized under section 505(k), including use of analytic tools and querying capabilities developed to implement the active postmarket surveillance system described in this section, public health activities as described in section 164.512(b)(1)(iii) of title 45, Code of Federal Regulations (or any successor regulation) and exempted as a public health activity as described in section 46.101(b)(5) of title 46, Code of Federal Regulations (or any successor regulation).

21 "(c) AUTHORIZATION OF APPROPRIATIONS.—There are 22 authorized to be appropriated to carry out this section 23 \$3,000,000 for each of fiscal years 2016 through 2020.".

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1 SEC. 2063. STREAMLINED DATA REVIEW PROGRAM.

- 2 (a) In General.—Chapter V of the Federal Food,
- 3 Drug, and Cosmetic Act, as amended by section 2062, is
- 4 further amended by inserting after section 505G of such Act
- 5 the following:

6 "SEC. 505H. STREAMLINED DATA REVIEW PROGRAM.

- 7 "(a) In General.—The Secretary shall establish a
- 8 streamlined data review program under which a holder of
- 9 an approved application submitted under section 505(b)(1)
- 10 or under section 351(a) of the Public Health Service Act
- 11 may, to support the approval or licensure (as applicable)
- 12 of the use of the drug that is the subject of such approved
- 13 application for a new qualified indication, submit qualified
- 14 data summaries.
- 15 "(b) Eligibility.—In carrying out the streamlined
- 16 data review program under subsection (a), the Secretary
- 17 may authorize the holder of the approved application to in-
- 18 clude one or more qualified data summaries described in
- 19 subsection (a) in a supplemental application if—
- 20 "(1) the drug has been approved under section
- 21 505(c) of this Act or licensed under section 351(a) of
- 22 the Public Health Service Act for one or more indica-
- 23 tions, and such approval or licensure remains in ef-
- 24 *fect*;
- 25 "(2) the supplemental application is for ap-
- 26 proval or licensure (as applicable) under such section

1	505(c) or 351(a) of the use of the drug for a new
2	qualified indication under such section 505(c) or
3	351(a);
4	"(3) there is an existing database acceptable to
5	the Secretary regarding the safety of the drug devel-
6	oped for one or more indications of the drug approved
7	under such section 505(c) or licensed under such sec-
8	tion 351(a);
9	"(4) the supplemental application incorporates
10	or supplements the data submitted in the application
11	for approval or licensure referred to in paragraph (1),
12	and
13	"(5) the full data sets used to develop the quali-
14	fied data summaries are submitted, unless the Sec-
15	retary determines that the full data sets are not re-
16	quired.
17	"(c) Public Availability of Information on Pro-
18	GRAM.—The Secretary shall post on the public website of
19	the Food and Drug Administration and update annually—
20	"(1) the number of applications reviewed under
21	the streamlined data review program;
22	"(2) the average time for completion of review
23	under the streamlined data review program versus
24	other review of applications for new indications: and

1	"(3) the number of applications reviewed under
2	the streamlined data review program for which the
3	Food and Drug Administration made use of full data
4	sets in addition to the qualified data summary.
5	"(d) Definitions.—In this section:
6	"(1) The term 'qualified indication' means—
7	"(A) an indication for the treatment of can-
8	cer, as determined appropriate by the Secretary;
9	or
10	"(B) such other types of indications as the
11	Secretary determines to be subject to the stream-
12	lined data review program under this section.
13	"(2) The term 'qualified data summary' means
14	a summary of clinical data intended to demonstrate
15	safety and effectiveness with respect to a qualified in-
16	dication for use of a drug.".
17	(b) Sense of Congress.—It is the sense of Congress
18	that the streamlined data review program under section
19	505H of the Federal Food, Drug, and Cosmetic Act, as
20	added by subsection (a), should enable the Food and Drug
21	Administration to make approval decisions for certain sup-
22	plemental applications based on qualified data summaries
23	(as defined in such section 505H).
24	(c) Guidance; Regulations.—The Commissioner of
25	Food and Druas—

1	(1) shall—
2	(A) issue final guidance for implementation
3	of the streamlined data review program estab-
4	lished under section 505H of the Federal Food,
5	Drug, and Cosmetic Act, as added by subsection
6	(a), not later than 24 months after the date of
7	enactment of this Act; and
8	(B) include in such guidance the process for
9	expanding the types of indications to be subject
10	to the streamlined data review program, as au-
11	thorized by section $505H(c)(1)(B)$ of such Act;
12	and
13	(2) in addition to issuing guidance under para-
14	graph (1), may issue such regulations as may be nec-
15	essary for implementation of the program.
16	Subtitle E—Expediting Patient
17	Access
18	SEC. 2081. SENSE OF CONGRESS.
19	It is the sense of Congress that the Food and Drug Ad-
20	ministration should continue to expedite the approval of
21	drugs designated as breakthrough therapies pursuant to sec-
22	tion 506(a) of the Federal Food, Drug, and Cosmetic Act
23	(21 U.S.C. 356(a)) by approving drugs so designated as
24	early as possible in the clinical development process, regard-
25	less of the phase of development, provided that the Secretary

- 1 of Health and Human Services determines that an applica-
- 2 tion for such a drug meets the standards of evidence of safe-
- 3 ty and effectiveness under section 505 of such Act (21 U.S.C.
- 4 355), including the substantial evidence standard under
- 5 subsection (d) of such section or under section 351(a) of the
- 6 Public Health Service Act (42 U.S.C. 262(a)).
- 7 SEC. 2082. EXPANDED ACCESS POLICY.
- 8 Chapter V of the Federal Food, Drug, and Cosmetic
- 9 Act is amended by inserting after section 561 (21 U.S.C.
- 10 360bbb) the following:
- 11 "SEC. 561A. EXPANDED ACCESS POLICY REQUIRED FOR IN-
- 12 **VESTIGATIONAL DRUGS.**
- 13 "(a) In General.—The manufacturer or distributor
- 14 of one or more investigational drugs for the diagnosis, mon-
- 15 itoring, or treatment of one or more serious diseases or con-
- 16 ditions shall make publicly available the policy of the man-
- 17 ufacturer or distributor on evaluating and responding to
- 18 requests submitted under section 561(b) for provision of
- 19 such a drug. A manufacturer or distributor may satisfy the
- 20 requirement of the preceding sentence by posting such policy
- 21 as generally applicable to all of such manufacturer's or dis-
- 22 tributor's investigational drugs.
- 23 "(b) Content of Policy.—A policy described in sub-
- 24 section (a) shall include making publicly available—

1	"(1) contact information for the manufacturer or
2	distributor to facilitate communication about requests
3	described in subsection (a);
4	"(2) procedures for making such requests;
5	"(3) the general criteria the manufacturer or dis-
6	tributor will consider or use to approve such requests;
7	and
8	"(4) the length of time the manufacturer or dis-
9	tributor anticipates will be necessary to acknowledge
10	receipt of such requests.
11	"(c) No Guarantee of Access.—The posting of poli-
12	cies by manufacturers and distributors under subsection (a)
13	shall not serve as a guarantee of access to any specific inves-
14	tigational drug by any individual patient.
15	"(d) Revised Policy.—A manufacturer or dis-
16	tributor that has made a policy publicly available as re-
17	quired by this section may revise the policy at any time.
18	"(e) Application.—This section shall apply to a
19	manufacturer or distributor with respect to an investiga-
20	tional drug beginning on the later of—
21	"(1) the date that is 60 days after the date of en-
22	actment of the 21st Century Cures Act; or
23	"(2) the first initiation of a phase 2 or phase 3
24	study (as such terms are defined in section 312.21(b)
25	and (c) of title 21, Code of Federal Regulations (or

1	any successor regulations)) with respect to such inves-
2	tigational new drug.".
3	SEC. 2083. FINALIZING DRAFT GUIDANCE ON EXPANDED
4	ACCESS.
5	(a) In General.—Not later than 12 months after the
6	date of enactment of this Act, the Secretary of Health and
7	Human Services shall finalize the draft guidance entitled
8	"Expanded Access to Investigational Drugs for Treatment
9	Use—Qs & As" and dated May 2013.
10	(b) Contents.—The final guidance referred to in sub-
11	section (a) shall clearly define how the Secretary of Health
12	and Human Services interprets and uses adverse drug event
13	data reported by investigators in the case of data reported
14	from use under a request submitted under section 561(b)
15	of the Federal Food, Drug, and Cosmetic Act (21 U.S.C.
16	360bbb(b)).
17	Subtitle F—Facilitating Respon-
18	sible Manufacturer Communica-
19	tions
20	SEC. 2101. FACILITATING DISSEMINATION OF HEALTH CARE
21	ECONOMIC INFORMATION.
22	Section 502(a) of the Federal Food, Drug, and Cos-
23	metic Act (21 U.S.C. 352(a)) is amended—
24	(1) by striking "(a) If its" and inserting "(a)(1)
25	If its":

- (2) by striking "a formulary committee, or other similar entity, in the course of the committee or the entity carrying out its responsibilities for the selection of drugs for managed care or other similar organizations" and inserting "a payor, formulary committee, or other similar entity with knowledge and expertise in the area of health care economic analysis, carrying out its responsibilities for the selection of drugs for coverage or reimbursement";
 - (3) by striking "directly relates" and inserting "relates":
 - (4) by striking "and is based on competent and reliable scientific evidence. The requirements set forth in section 505(a) or in section 351(a) of the Public Health Service Act shall not apply to health care economic information provided to such a committee or entity in accordance with this paragraph" and inserting ", is based on competent and reliable scientific evidence, and includes, where applicable, a conspicuous and prominent statement describing any material differences between the health care economic information and the labeling approved for the drug under section 505 or under section 351 of the Public Health Service Act. The requirements set forth in section 505(a) or in subsections (a) and (k) of section

1	351 of the Public Health Service Act shall not apply
2	to health care economic information provided to such
3	a payor, committee, or entity in accordance with this
4	paragraph"; and
5	(5) by striking "In this paragraph, the term"
6	and all that follows and inserting the following:
7	"(2)(A) For purposes of this paragraph, the term
8	health care economic information' means any analysis (in-
9	cluding the clinical data, inputs, clinical or other assump-
10	tions, methods, results, and other components underlying or
11	comprising the analysis) that identifies, measures, or de-
12	scribes the economic consequences, which may be based on
13	the separate or aggregated clinical consequences of the rep-
14	resented health outcomes, of the use of a drug. Such analysis
15	may be comparative to the use of another drug, to another
16	health care intervention, or to no intervention.
17	"(B) Such term does not include any analysis that re-
18	lates only to an indication that is not approved under sec-
19	tion 505 or under section 351 of the Public Health Service
20	Act for such drug.".
21	SEC. 2102. FACILITATING RESPONSIBLE COMMUNICATION
22	OF SCIENTIFIC AND MEDICAL DEVELOP-
23	MENTS.
24	(a) GUIDANCE.—Not later than 18 months after the
25	date of enactment of this Act, the Secretary of Health and

- 1 Human Services shall issue draft guidance on facilitating
- 2 the responsible dissemination of truthful and nonmisleading
- 3 scientific and medical information not included in the ap-
- 4 proved labeling of drugs and devices.
- 5 (b) Definition.—In this section, the terms "drug"
- 6 and "device" have the meaning given to such terms in sec-
- 7 tion 201 of the Federal Food, Drug, and Cosmetic Act (21
- 8 U.S.C. 321).

9 Subtitle G—Antibiotic Drug

10 **Development**

- 11 SEC. 2121. APPROVAL OF CERTAIN DRUGS FOR USE IN A
- 12 LIMITED POPULATION OF PATIENTS.
- 13 (a) Purpose.—The purpose of this section is to help
- 14 to expedite the development and availability of treatments
- 15 for serious or life-threatening bacterial or fungal infections
- 16 in patients with unmet needs, while maintaining safety and
- 17 effectiveness standards for such treatments, taking into ac-
- 18 count the severity of the infection and the availability or
- 19 lack of alternative treatments.
- 20 (b) Approval of Certain Antibacterial and
- 21 Antifungal Drugs.—Section 505 of the Federal Food,
- 22 Drug, and Cosmetic Act (21 U.S.C. 355), as amended by
- 23 section 2001, is further amended by adding at the end the
- 24 following new subsection:

1	"(z) Approval of Certain Antibacterial and
2	Antifungal Drugs for Use in a Limited Population
3	OF PATIENTS.—
4	"(1) Process.—At the request of the sponsor of
5	an antibacterial or antifungal drug that is intended
6	to treat a serious or life-threatening infection, the
7	Secretary—
8	"(A) may execute a written agreement with
9	the sponsor on the process for developing data to
10	support an application for approval of such
11	drug, for use in a limited population of patients
12	in accordance with this subsection;
13	"(B) shall proceed in accordance with this
14	subsection only if a written agreement is reached
15	$under\ subparagraph\ (A);$
16	"(C) shall provide the sponsor with an op-
17	portunity to request meetings under paragraph
18	(2);
19	"(D) if a written agreement is reached
20	under subparagraph (A), may approve the drug
21	under this subsection for such use—
22	"(i) in a limited population of pa-
23	tients for which there is an unmet medical
24	need;

1	"(ii) based on a streamlined develop-
2	ment program; and
3	"(iii) only if the standards for ap-
4	proval under subsections (c) and (d) of this
5	section or licensure under section 351 of the
6	Public Health Service Act, as applicable,
7	are met; and
8	"(E) in approving a drug in accordance
9	with this subsection, subject to subparagraph
10	(D)(iii), may rely upon—
11	"(i) traditional endpoints, alternate
12	endpoints, or a combination of traditional
13	and alternate endpoints, and, as appro-
14	priate, data sets of a limited size; and
15	$``(ii)(I)\ additional\ data,\ including\ pre-$
16	clinical, pharmacologic, or pathophysiologic
17	evidence;
18	"(II) nonclinical susceptibility and
19	$pharmacokinetic\ data;$
20	"(III) data from phase 2 clinical
21	trials; and
22	"(IV) such other confirmatory evidence
23	as the Secretary determines appropriate to
24	approve the drug.
25	"(2) Formal meetings.—

1	"(A) In general.—To help to expedite and
2	facilitate the development and review of a drug
3	for which a sponsor intends to request approval
4	in accordance with this subsection, the Secretary
5	may, at the request of the sponsor, conduct meet-
6	ings that provide early consultation, timely ad-
7	vice, and sufficient opportunities to develop an
8	agreement described in paragraph (1)(A) and
9	help the sponsor design and conduct a drug de-
10	velopment program as efficiently as possible, in-
11	cluding the following types of meetings:
12	"(i) An early consultation meeting.
13	"(ii) An assessment meeting.
14	$``(iii)\ A\ postapproval\ meeting.$
15	"(B) No altering of goals.—Nothing in
16	this paragraph shall be construed to alter agreed
17	upon goals and procedures identified in the let-
18	ters described in section 101(b) of the Prescrip-
19	tion Drug User Fee Amendments of 2012.
20	"(C) Breakthrough therapies.—In the
21	case of a drug designated as a breakthrough ther-
22	apy under section 506(a), the sponsor of such
23	drug may elect to utilize meetings provided
24	under such section with respect to such drug in
25	lieu of meetings described in subparagraph (A).

- "(3) Labeling requirement.—The labeling of an antibacterial or antifungal drug approved in accordance with this subsection shall contain the statement 'Limited Population' in a prominent manner and adjacent to, and not more prominent than, the brand name of the product. The prescribing information for such antibacterial or antifungal drug required by section 201.57 of title 21, Code of Federal Regulations (or any successor regulation) shall also include the following statement: 'This drug is indicated for use in a limited and specific population of patients.'
 - "(4) Promotional materials.—The provisions of section 506(c)(2)(B) shall apply with respect to approval in accordance with this subsection to the same extent and in the same manner as such provisions apply with respect to accelerated approval in accordance with section 506(c)(1).
 - "(5) TERMINATION OF REQUIREMENTS OR CON-DITIONS.—If a drug is approved in accordance with this subsection for an indication in a limited population of patients and is subsequently approved or licensed under this section or section 351 of the Public Health Service Act, other than in accordance with this subsection, for—

1	"(A) the same indication and the same con-
2	ditions of use, the Secretary shall remove any la-
3	beling requirements or postmarketing conditions
4	that were made applicable to the drug under this
5	$subsection;\ or$
6	"(B) a different indication or condition of
7	use, the Secretary shall not apply the labeling re-
8	quirements and postmarketing conditions that
9	were made applicable to the drug under this sub-
10	section to the subsequent approval of the drug for
11	such different indication or condition of use.
12	"(6) Relation to other provisions.—Nothing
13	in this subsection shall be construed to prohibit the
14	approval of a drug for use in a limited population
15	of patients in accordance with this subsection, in
16	combination with—
17	"(A) an agreement on the design and size of
18	a clinical trial pursuant to subparagraphs (B)
19	and (C) of subsection $(b)(5)$;
20	"(B) designation and treatment of the drug
21	as a breakthrough therapy under section 506(a);
22	"(C) designation and treatment of the drug
23	as a fast track product under section 506(b); or
24	"(D) accelerated approval of the drug in ac-
25	cordance with section $506(c)$.

1	"(7) Rule of construction.—Nothing in this
2	subsection shall be construed—
3	"(A) to alter the standards of evidence
4	under subsection (c) or (d) (including the sub-
5	stantial evidence standard in subsection (d));
6	"(B) to waive or otherwise preclude the ap-
7	plication of requirements under subsection (o);
8	"(C) to otherwise, in any way, limit the au-
9	thority of the Secretary to approve products pur-
10	suant to this Act and the Public Health Service
11	Act as authorized prior to the date of enactment
12	of this subsection; or
13	"(D) to restrict in any manner, the pre-
14	scribing of antibiotics or other products by
15	health care providers, or to otherwise limit or re-
16	strict the practice of health care.
17	"(8) Effective immediately.—The Secretary
18	shall have the authorities vested in the Secretary by
19	this subsection beginning on the date of enactment of
20	this subsection, irrespective of when and whether the
21	Secretary promulgates final regulations or guidance.
22	"(9) Definitions.—In this subsection:
23	"(A) Early consultation meeting.—The
24	term 'early consultation meeting' means a pre-

1	investigational new drug meeting or an end-of-
2	phase-1 meeting that—
3	"(i) is conducted to review and reach
4	a written agreement—
5	"(I) on the scope of the stream-
6	lined development plan for a drug for
7	which a sponsor intends to request ap-
8	proval in accordance with this sub-
9	section; and
10	"(II) which, as appropriate, may
11	include agreement on the design and
12	size of necessary preclinical and clin-
13	ical studies early in the development
14	process, including clinical trials whose
15	data are intended to form the primary
16	basis for an effectiveness claim; and
17	"(ii) provides an opportunity to dis-
18	cuss expectations of the Secretary regarding
19	studies or other information that the Sec-
20	retary deems appropriate for purposes of
21	applying paragraph (5), relating to the ter-
22	mination of labeling requirements or post-
23	marketing conditions.
24	"(B) Assessment meeting.—The term 'as-
25	sessment meeting' means an end-of-phase 2 meet-

ing, pre-new drug application meeting, or prebiologics license application meeting conducted to resolve questions and issues raised during the course of clinical investigations, and details addressed in the written agreement regarding postapproval commitments or expansion of approved uses.

- "(C) Postapproval meeting' means a meeting following initial approval or licensure of the drug for use in a limited population, to discuss any issues identified by the Secretary or the sponsor regarding postapproval commitments or expansion of approved uses.".
- 15 (c) GUIDANCE.—Not later than 18 months after the date of enactment of this Act, the Secretary of Health and 16 Human Services, acting through the Commissioner of Food and Drugs, shall issue draft guidance describing criteria, 18 process, and other general considerations for demonstrating 19 the safety and effectiveness of antibacterial and antifungal 20 21 drugs to be approved for use in a limited population in accordance with section 505(z) of the Federal Food, Drug, 23 and Cosmetic Act, as added by subsection (b).
- 24 (d) Conforming Amendments.—

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1	(1) Licensure of certain biological prod-
2	UCTS.—Section 351(j) of the Public Health Service
3	Act (42 U.S.C. 262(j)) is amended—
4	(A) by striking "(j)" and inserting "(j)(1)";
5	(B) by inserting "505(z)," after "505(p),";
6	and
7	(C) by adding at the end the following new
8	paragraph:
9	"(2) In applying section 505(z) of the Federal Food,
10	Drug, and Cosmetic Act to the licensure of biological prod-
11	ucts under this section—
12	"(A) references to an antibacterial or antifungal
13	drug that is intended to treat a serious or life-threat-
14	ening infection shall be construed to refer to a biologi-
15	cal product intended to treat a serious or life-threat-
16	ening bacterial or fungal infection; and
17	"(B) references to approval of a drug under sec-
18	tion 505(c) of such Act shall be construed to refer to
19	a licensure of a biological product under subsection
20	(a) of this section.".
21	(2) Misbranding.—Section 502 of the Federal
22	Food, Drug, and Cosmetic Act (21 U.S.C. 352) is
23	amended by adding at the end the following new sub-
24	section:

1 "(dd) If it is a drug approved in accordance with sec-

tion 505(z) and its labeling does not meet the requirements

3 under paragraph (3) of such subsection, subject to para-

4 graph (5) of such subsection.".

(e) Evaluation.—

- (1) Assessment.—Not later than 48 months after the date of enactment of this Act, the Secretary of Health and Human Services shall publish for public comment an assessment of the program established under section 505(z) of the Federal Food, Drug, and Cosmetic Act, as added by subsection (b). Such assessment shall determine if the limited-use pathway established under such section 505(z) has improved or is likely to improve patient access to novel antibacterial or antifungal treatments and assess how the pathway could be expanded to cover products for serious or life-threatening diseases or conditions beyond bacterial and fungal infections.
- (2) MEETING.—Not later than 90 days after the date of the publication of such assessment, the Secretary, acting through the Commissioner of Food and Drugs, shall hold a public meeting to discuss the findings of the assessment, during which public stakeholders may present their views on the success of the program established under section 505(z) of the Fed-

- 1 eral Food, Drug, and Cosmetic Act, as added by sub-
- 2 section (b), and the appropriateness of expanding
- 3 such program.
- 4 (f) Expansion of Program.—If the Secretary of
- 5 Health and Human Services determines, based on the as-
- 6 sessment under subsection (e)(1), evaluation of the assess-
- 7 ment, and any other relevant information, that the public
- 8 health would benefit from expansion of the limited-use path-
- 9 way established under section 505(z) of the Federal Food,
- 10 Drug, and Cosmetic Act (as added by subsection (b)) beyond
- 11 the drugs approved in accordance with such section, the
- 12 Secretary may expand such limited-use pathway in accord-
- 13 ance with such a determination. The approval of any drugs
- 14 under any such expansion shall be subject to the consider-
- 15 ations and requirements described in such section 505(z)
- 16 for purposes of expansion to other serious or life-threatening
- 17 diseases or conditions.
- 18 (g) Monitoring.—The Public Health Service Act is
- 19 amended by inserting after section 317T (42 U.S.C. 247b-
- 20 22) the following:
- 21 "SEC. 317U. MONITORING ANTIBACTERIAL AND
- 22 ANTIFUNGAL DRUG USE AND RESISTANCE.
- 23 "(a) Monitoring.—The Secretary shall use an appro-
- 24 priate monitoring system to monitor—

1	"(1) the use of antibacterial and antifungal
2	drugs, including those receiving approval or licensure
3	for a limited population pursuant to section 505(z) of
4	the Federal Food, Drug, and Cosmetic Act; and
5	"(2) changes in bacterial and fungal resistance
6	$to\ drugs.$
7	"(b) Public Availability of Data.—The Secretary
8	shall make summaries of the data derived from monitoring
9	under this section publicly available for the purposes of—
10	"(1) improving the monitoring of important
11	trends in antibacterial and antifungal resistance; and
12	"(2) ensuring appropriate stewardship of anti-
13	bacterial and antifungal drugs, including those re-
14	ceiving approval or licensure for a limited population
15	pursuant to section 505(z) of the Federal Food, Drug,
16	and Cosmetic Act.".
17	SEC. 2122. SUSCEPTIBILITY TEST INTERPRETIVE CRITERIA
18	FOR MICROORGANISMS.
19	(a) In General.—Section 511 of the Federal Food,
20	Drug, and Cosmetic Act (21 U.S.C. 360a) is amended to
21	read as follows:
22	"SEC. 511. IDENTIFYING AND UPDATING SUSCEPTIBILITY
23	TEST INTERPRETIVE CRITERIA FOR MICRO-
24	ORGANISMS.
25	"(a) Purpose: Identification of Criteria.—

1	"(1) Purpose.—The purpose of this section is to
2	provide the Secretary with an expedited, flexible
3	method for—
4	"(A) clearance or premarket approval of
5	antimicrobial susceptibility testing devices uti-
6	lizing updated, recognized susceptibility test in-
7	terpretive criteria to characterize the in vitro
8	susceptibility of particular bacteria, fungi, or
9	other microorganisms to antimicrobial drugs;
10	and
11	"(B) providing public notice of the avail-
12	ability of recognized interpretive criteria to meet
13	premarket submission requirements or other re-
14	quirements under this Act for antimicrobial sus-
15	ceptibility testing devices.
16	"(2) In general.—The Secretary shall identify
17	appropriate susceptibility test interpretive criteria
18	with respect to antimicrobial drugs—
19	"(A) if such criteria are available on the
20	date of approval of the drug under section 505
21	of this Act or licensure of the drug under section
22	351 of the Public Health Service Act (as applica-
23	ble), upon such approval or licensure; or

1	"(B) if such criteria are unavailable on
2	such date, on the date on which such criteria are
3	available for such drug.
4	"(3) Bases for initial identification.—The
5	Secretary shall identify appropriate susceptibility test
6	interpretive criteria under paragraph (2), based on
7	the Secretary's review of, to the extent available and
8	relevant—
9	"(A) preclinical and clinical data, includ-
10	ing pharmacokinetic, pharmacodynamic, and ep-
11	$idemiological\ data;$
12	"(B) Bayesian and pharmacometric statis-
13	tical methodologies; and
14	"(C) such other evidence and information as
15	the Secretary considers appropriate.
16	"(b) Susceptibility Test Interpretive Criteria
17	Website.—
18	"(1) In General.—Not later than 1 year after
19	the date of the enactment of the 21st Century Cures
20	Act, the Secretary shall establish, and maintain there-
21	after, on the website of the Food and Drug Adminis-
22	tration, a dedicated website that contains a list of
23	any appropriate new or updated susceptibility test
24	interpretive criteria standards in accordance with

1	paragraph (2) (referred to in this section as the In-
2	terpretive Criteria Website').
3	"(2) Listing of susceptibility test inter-
4	PRETIVE CRITERIA STANDARDS.—
5	"(A) In general.—The list described in
6	paragraph (1) shall consist of any new or up-
7	dated susceptibility test interpretive criteria
8	standards that are—
9	"(i) established by a nationally or
10	internationally recognized standard devel-
11	opment organization that—
12	"(I) establishes and maintains
13	procedures to address potential con-
14	flicts of interest and ensure trans-
15	$parent\ decision making;$
16	"(II) holds open meetings to en-
17	sure that there is an opportunity for
18	public input by interested parties, and
19	establishes and maintains processes to
20	ensure that such input is considered in
21	$decision making;\ and$
22	"(III) permits its standards to be
23	made publicly available, through the
24	National Library of Medicine or an-

1	other similar source acceptable to the
2	Secretary; and
3	"(ii) recognized in whole, or in part,
4	by the Secretary under subsection (c).
5	"(B) Other list.—The Interpretive Cri-
6	teria Website shall, in addition to the list de-
7	scribed in subparagraph (A), include a list of in-
8	terpretive criteria, if any, that the Secretary has
9	determined to be appropriate with respect to le-
10	gally marketed antimicrobial drugs, where—
11	"(i) the Secretary does not recognize,
12	in whole or in part, an interpretive criteria
13	standard described under subparagraph (A)
14	otherwise applicable to such a drug;
15	"(ii) the Secretary withdraws under
16	subsection $(c)(1)(B)$ recognition of a stand-
17	ard, in whole or in part, otherwise applica-
18	ble to such a drug;
19	"(iii) the Secretary approves an appli-
20	cation under section 505 of this Act or sec-
21	tion 351 of the Public Health Service Act,
22	as applicable, with respect to marketing of
23	such a drug for which there are no relevant
24	interpretive criteria included in a standard

1	recognized by the Secretary under sub-
2	section (c); or
3	"(iv) because the characteristics of such
4	a drug differ from other drugs with the
5	same active ingredient, the interpretive cri-
6	teria with respect to such drug—
7	"(I) differ from otherwise applica-
8	ble interpretive criteria included in a
9	standard listed under subparagraph
10	(A) or interpretive criteria otherwise
11	listed under this subparagraph; and
12	"(II) are determined by the Sec-
13	retary to be appropriate for the drug.
14	"(C) REQUIRED STATEMENTS OF LIMITA-
15	Tions of information.—The Interpretive Cri-
16	teria Website shall include the following:
17	"(i) A statement that—
18	"(I) the website provides informa-
19	tion about the susceptibility of bac-
20	teria, fungi, or other microorganisms
21	to a certain drug (or drugs); and
22	"(II) the safety and efficacy of the
23	drug in treating clinical infections due
24	to such bacteria, fungi, or other micro-
25	organisms may not have been estab-

1	lished in adequate and well-controlled
2	clinical trials and the clinical signifi-
3	cance of such susceptibility informa-
4	tion in such trials is unknown.
5	"(ii) A statement that directs health
6	care practitioners to consult the approved
7	product labeling for specific drugs to deter-
8	mine the uses for which the Food and Drug
9	Administration has approved the product.
10	"(iii) Any other statement that the
11	Secretary determines appropriate to ade-
12	quately convey the limitations of the data
13	supporting susceptibility test interpretive
14	criteria standard listed on the website.
15	"(3) Notice.—Not later than the date on which
16	the Interpretive Criteria Website is established, the
17	Secretary shall publish a notice of that establishment
18	in the Federal Register.
19	"(4) Inapplicability of misbranding provi-
20	SION.—The inclusion in the approved labeling of an
21	antimicrobial drug of a reference or hyperlink to the
22	Interpretive Criteria Website, in and of itself, shall
23	not cause the drug to be misbranded in violation of
24	section 502, or the regulations promulgated there-
25	under.

1	"(5) Trade secrets and confidential infor-
2	MATION.—Nothing in this section shall be construed
3	as authorizing the Secretary to disclose any informa-
4	tion that is a trade secret or confidential information
5	subject to section 552(b)(4) of title 5, United States
6	Code.
7	"(c) Recognition of Susceptibility Test Inter-
8	PRETIVE CRITERIA FROM STANDARD DEVELOPMENT ORGA-
9	NIZATIONS.—
10	"(1) In general.—Beginning on the date of the
11	establishment of the Interpretive Criteria Website, and
12	at least every 6 months thereafter, the Secretary
13	shall—
14	"(A) evaluate any appropriate new or up-
15	dated susceptibility test interpretive criteria
16	standards established by a nationally or inter-
17	nationally recognized standard development or-
18	$ganization \ described \ in \ subsection \ (b)(2)(A)(i);$
19	and
20	"(B) publish on the public website of the
21	Food and Drug Administration a notice—
22	"(i) withdrawing recognition of any
23	different susceptibility test interpretive cri-
24	teria standard, in whole or in part:

1	"(ii) recognizing the new or updated
2	standards;
3	"(iii) recognizing one or more parts of
4	the new or updated interpretive criteria
5	specified in such a standard and declining
6	to recognize the remainder of such standard;
7	and
8	"(iv) making any necessary updates to
9	the lists under subsection $(b)(2)$.
10	"(2) Bases for updating interpretive cri-
11	TERIA STANDARDS.—In evaluating new or updated
12	susceptibility test interpretive criteria standards
13	under paragraph (1)(A), the Secretary may con-
14	sider—
15	"(A) the Secretary's determination that
16	such a standard is not applicable to a particular
17	drug because the characteristics of the drug differ
18	from other drugs with the same active ingredient;
19	"(B) information provided by interested
20	third parties, including public comment on the
21	annual compilation of notices published under
22	paragraph (3);
23	"(C) any bases used to identify suscepti-
24	bility test interpretive criteria under subsection
25	(a)(2); and

1	"(D) such other information or factors as
2	the Secretary determines appropriate.
3	"(3) Annual compilation of notices.—Each
4	year, the Secretary shall compile the notices published
5	under paragraph (1)(B) and publish such compila-
6	tion in the Federal Register and provide for public
7	comment. If the Secretary receives comments, the Sec-
8	retary will review such comments and, if the Sec-
9	retary determines appropriate, update pursuant to
10	this subsection susceptibility test interpretive criteria
11	standards—
12	"(A) recognized by the Secretary under this
13	subsection; or
14	"(B) otherwise listed on the Interpretive
15	Criteria Website under subsection (b)(2).
16	"(4) Relation to Section 514(c).—Any suscep-
17	tibility test interpretive standard recognized under
18	this subsection or any criteria otherwise listed under
19	subsection $(b)(2)(B)$ shall be deemed to be recognized
20	as a standard by the Secretary under section
21	514(c)(1).
22	"(5) Voluntary use of interpretive cri-
23	TERIA.—Nothing in this section prohibits a person
24	from seeking approval or clearance of a drug or de-
25	vice, or changes to the drug or the device, on the basis

1	of susceptibility test interpretive criteria standards
2	which differ from those recognized pursuant to para-
3	graph(1).
4	"(d) Antimicrobial Drug Labeling.—
5	"(1) Drugs marketed prior to establish-
6	MENT OF INTERPRETIVE CRITERIA WEBSITE.—With
7	respect to an antimicrobial drug lawfully introduced
8	or delivered for introduction into interstate commerce
9	for commercial distribution before the establishment of
10	the Interpretive Criteria Website, a holder of an ap-
11	proved application under section 505 of this Act or
12	section 351 of the Public Health Service Act, as ap-
13	plicable, for each such drug—
14	"(A) not later than 1 year after establish
15	ment of the Interpretive Criteria Website, shall
16	submit to the Secretary a supplemental applica-
17	tion for purposes of changing the drug's labeling
18	to substitute a reference or hyperlink to such
19	Website for any susceptibility test interpretive
20	criteria and related information; and
21	"(B) may begin distribution of the drug in
22	volved upon receipt by the Secretary of the sup-
23	plemental application for such change.
24	"(2) Drugs marketed subsequent to estab-
25	LISHMENT OF INTEDDDETINE COURTLA WEDSITE

1	With respect to antimicrobial drugs lawfully intro-
2	duced or delivered for introduction into interstate
3	commerce for commercial distribution on or after the
4	date of the establishment of the Interpretive Criteria
5	Website, the labeling for such a drug shall include, in
6	lieu of susceptibility test interpretive criteria and re-
7	lated information, a reference to such Website.
8	"(e) Special Condition for Marketing of Anti-
9	MICROBIAL SUSCEPTIBILITY TESTING DEVICES.—
10	"(1) In General.—Notwithstanding sections
11	501, 502, 510, 513, and 515, if the conditions speci-
12	fied in paragraph (2) are met (in addition to other
13	applicable provisions under this chapter) with respect
14	to an antimicrobial susceptibility testing device de-
15	scribed in subsection (f)(1), the Secretary may au-
16	thorize the marketing of such device for a use de-
17	scribed in such subsection.
18	"(2) Conditions applicable to anti-
19	MICROBIAL SUSCEPTIBILITY TESTING DEVICES.—The
20	conditions specified in this paragraph are the fol-
21	lowing:
22	"(A) The device is used to make a deter-
23	mination of susceptibility using susceptibility
24	test interpretive criteria that are—

1	"(i) included in a standard recognized
2	by the Secretary under subsection (c); or
3	"(ii) otherwise listed on the Interpre-
4	tive Criteria Website under subsection
5	(b)(2).
6	"(B) The labeling of such device promi-
7	nently and conspicuously—
8	"(i) includes a statement that—
9	"(I) the device provides informa-
10	tion about the susceptibility of bacteria
11	and fungi to certain drugs; and
12	"(II) the safety and efficacy of
13	such drugs in treating clinical infec-
14	tions due to such bacteria or fungi
15	may not have been established in ade-
16	quate and well-controlled clinical trials
17	and the clinical significance of such
18	susceptibility information in those in-
19	stances is unknown;
20	"(ii) includes a statement directing
21	health care practitioners to consult the ap-
22	proved labeling for drugs tested using such
23	a device, to determine the uses for which the
24	Food and Drug Administration has ap-
25	proved such drugs; and

1	"(iii) includes any other statement the
2	Secretary determines appropriate to ade-
3	quately convey the limitations of the data
4	supporting the interpretive criteria de-
5	scribed in subparagraph (A).
6	"(f) Definitions.—In this section:
7	"(1) The term 'antimicrobial susceptibility test-
8	ing device' means a device that utilizes susceptibility
9	test interpretive criteria to determine and report the
10	in vitro susceptibility of certain microorganisms to a
11	drug (or drugs).
12	"(2) The term 'qualified infectious disease prod-
13	uct' means a qualified infectious disease product des-
14	$ignated\ under\ section\ 505 E(d).$
15	"(3) The term 'susceptibility test interpretive cri-
16	teria' means—
17	"(A) one or more specific numerical values
18	which characterize the susceptibility of bacteria
19	or other microorganisms to the drug tested; and
20	"(B) related categorizations of such suscep-
21	tibility, including categorization of the drug as
22	susceptible, intermediate, resistant, or such other
23	term as the Secretary determines appropriate.

1	"(4)(A) The term 'antimicrobial drug' means,
2	subject to subparagraph (B), a systemic antibacterial
3	or antifungal drug that—
4	"(i) is intended for human use in the treat-
5	ment of a disease or condition caused by a bac-
6	terium or fungus;
7	"(ii) may include a qualified infectious dis-
8	ease product designated under section $505E(d)$;
9	and
10	"(iii) is subject to section 503(b)(1).
11	"(B) If provided by the Secretary through regu-
12	lations, such term may include—
13	"(i) drugs other than systemic antibacterial
14	and antifungal drugs; and
15	"(ii) biological products (as such term is de-
16	fined in section 351 of the Public Health Service
17	Act) to the extent such products exhibit anti-
18	$microbial\ activity.$
19	"(g) Rule of Construction.—Nothing in this sec-
20	tion shall be construed—
21	"(1) to alter the standards of evidence—
22	"(A) under subsection (c) or (d) of section
23	505, including the substantial evidence standard
24	in section $505(d)$, or under section 351 of the
25	Public Health Service Act (as applicable); or

1	"(B) with respect to marketing authoriza-
2	tion for devices, under section 510, 513, or 515;
3	"(2) to apply with respect to any drug, device,
4	or biological product, in any context other than—
5	"(A) an antimicrobial drug; or
6	"(B) an antimicrobial susceptibility testing
7	device that uses susceptibility test interpretive
8	criteria to characterize and report the in vitro
9	susceptibility of certain bacteria, fungi, or other
10	microorganisms to antimicrobial drugs in ac-
11	cordance with this section; or
12	"(3) unless specifically stated, to have any effect
13	on authorities provided under other sections of this
14	Act, including any regulations issued under such sec-
15	tions.".
16	(b) Conforming Amendments.—
17	(1) Repeal of related authority.—Section
18	1111 of the Food and Drug Administration Amend-
19	ments Act of 2007 (42 U.S.C. 247d-5a; relating to
20	identification of clinically susceptible concentrations
21	of antimicrobials) is repealed.
22	(2) Clerical amendment.—The table of con-
23	tents in section 2 of the Food and Drug Administra-
24	tion Amendments Act of 2007 is amended by striking
25	the item relating to section 1111.

1 (3) Misbranding.—Section 502 of the Federal 2 Food, Drug, and Cosmetic Act (21 U.S.C. 352), as 3 amended by section 2121, is further amended by add-4 ing at the end the following: 5 "(ee) If it is an antimicrobial drug and its labeling fails to conform with the requirements under section 7 511(d).". 8 (4) Recognition of interpretive criteria as 9 DEVICE STANDARD.—Section 514(c)(1)(A) of the Fed-10 eral Food, Drug, and Cosmetic Act (21 U.S.C. 11 360d(c)(1)(A)) is amended by inserting after "the 12 Secretary shall, by publication in the Federal Reg-13 ister" the following: "(or, with respect to suscepti-14 bility test interpretive criteria or standards recog-15 nized or otherwise listed under section 511, by posting 16 on the Interpretive Criteria Website in accordance 17 with such section)". 18 (c) Report to Congress.—Not later than two years after the date of enactment of this Act, the Secretary of 19 Health and Human Services shall submit to the Committee 21 on Energy and Commerce of the House of Representatives and the Committee on Health, Education, Labor and Pensions of the Senate a report on the progress made in implementing section 511 of the Federal Food, Drug, and Cos-

metic Act (21 U.S.C. 360a), as amended by this section.

1	(d) Requests for Updates to Interpretive Cri-
2	TERIA WEBSITE.—Chapter 35 of title 44, United States
3	Code, shall not apply to the collection of information from
4	interested parties regarding the updating of lists under
5	paragraph (2) of subsection (b) section 511 of the Federal
6	Food, Drug, and Cosmetic Act (as amended by subsection
7	(a)) and posted on the Interpretive Criteria Website estab-
8	lished under paragraph (1) of such subsection (b).
9	(e) No Effect on Health Care Practice.—Noth-
10	ing in this subtitle (including the amendments made by this
11	subtitle) shall be construed to restrict, in any manner, the
12	prescribing or administering of antibiotics or other prod-
13	ucts by health care practitioners, or to limit the practice
14	of health care.
15	SEC. 2123. ENCOURAGING THE DEVELOPMENT AND USE OF
16	NEW ANTIMICROBIAL DRUGS.
17	(a) Additional Payment for New Antimicrobial
18	Drugs Under Medicare.—
19	(1) In General.—Section 1886(d)(5) of the So-
20	cial Security Act (42 U.S.C. $1395ww(d)(5)$) is
21	amended by adding at the end the following new sub-

"(M)(i) As part of the annual rulemaking under this
subsection for payment for subsection (d) hospitals for each

paragraph:

22

1	fiscal year beginning with fiscal year 2018, the Secretary
2	shall—
3	"(I) include publication of a list of the new anti-
4	microbial drugs for such fiscal year; and
5	"(II) with respect to discharges by eligible hos-
6	pitals that involve a drug so published, provide for an
7	additional payment to be made under this subsection
8	in accordance with the provisions of this subpara-
9	graph.
10	"(ii) Additional payments may not be made for a drug
11	under this subparagraph—
12	"(I) other than during the 5-fiscal-year period
13	beginning with the fiscal year for which the drug is
14	first included in the publication described in clause
15	(i)(I); and
16	"(II) with respect to which payment has even
17	been made pursuant to subparagraph (K).
18	"(iii) For purposes of this subparagraph, the term
19	'new antimicrobial drug' means a product that is approved
20	for use, or a product for which an indication is first ap-
21	proved for use, by the Food and Drug Administration on
22	or after December 1, 2014, and that the Food and Drug
23	Administration determines—
24	"(I) either—

1	"(aa) is intended to treat an infection
2	caused by, or likely to be caused by, a qualifying
3	pathogen (as defined under section $505E(f)$ of the
4	Federal Food, Drug, and Cosmetic Act); or
5	"(bb) meets the definition of a qualified in-
6	fectious disease product under section $505E(g)$ of
7	the Federal Food, Drug, and Cosmetic Act; and
8	"(II) is intended to treat an infection—
9	"(aa) for which there is an unmet medical
10	need; and
11	"(bb) which is associated with high rates of
12	mortality or significant patient morbidity, as
13	determined in consultation with the Director of
14	the Centers for Disease Control and Prevention
15	and the infectious disease professional commu-
16	nity.
17	Such determination may be revoked only upon a finding
18	that the request for such determination contained an untrue
19	statement of material fact.
20	"(iv) For purposes of this subparagraph, the term 'eli-
21	gible hospital' means a subsection (d) hospital that partici-
22	pates in the National Healthcare Safety Network of the Cen-
23	ters for Disease Control and Prevention (or, to the extent
24	a similar surveillance system reporting program that in-
25	cludes reporting about antimicrobial drugs is determined

- 1 by the Secretary to be available to such hospitals, such simi-
- 2 lar surveillance system as the Secretary may specify).
- 3 "(v)(I) Subject to the succeeding provisions of this
- 4 clause, the additional payment under this subparagraph,
- 5 with respect to a drug, shall be in the amount provided
- 6 for such drug under section 1847A.
- 7 "(II) The Secretary shall, as part of the rulemaking
- 8 referred to in clause (i) for each fiscal year, estimate—
- 9 "(aa) the total amount of the additional pay-
- 10 ments that will be made under this subsection pursu-
- ant to this subparagraph for discharges in such fiscal
- 12 year without regard to the application of subclause
- 13 (III); and
- 14 "(bb) the total program payments to be made
- 15 under this subsection for all discharges in such fiscal
- 16 year.
- 17 "(III) If the estimated total amount described in sub-
- 18 clause (II)(aa) for a fiscal year exceeds the applicable per-
- 19 centage of the estimated total program payments described
- 20 in subclause (II)(bb) for such fiscal year, the Secretary shall
- 21 reduce in a pro rata manner the amount of each additional
- 22 payment under this subsection pursuant to this subpara-
- 23 graph for such fiscal year in order to ensure that the total
- 24 amount of the additional payments under this subsection
- 25 pursuant to this subparagraph for such fiscal year do not

1	exceed the applicable percentage of the estimated total pro-
2	gram payments described in subclause (II)(bb) for such fis-
3	cal year.
4	"(IV) For purposes of subclause (III), the term 'appli-
5	cable percentage' means 0.03 percent.".
6	(2) Conforming amendments.—
7	(A) NO DUPLICATIVE NTAP PAYMENTS.—
8	Section $1886(d)(5)(K)(vi)$ of the Social Security
9	Act (42 U.S.C. $1395ww(d)(5)(K)(vi)$) is amended
10	by inserting "if additional payment has never
11	been made under this subsection pursuant to sub-
12	paragraph (M) with respect to the service or
13	technology" after "if the service or technology".
14	(B) Access to price information.—Sec-
15	tion 1927(b)(3)(A)(iii) of the Social Security Act
16	(42 U.S.C. 1396r-8(b)(3)(A)(iii)) is amended—
17	(i) in subclause (II), by inserting "or
18	under section 1886(d) pursuant to para-
19	graph (5)(M) of such section," after
20	"1847A,"; and
21	(ii) in the matter following subclause
22	(III), by inserting "or section
23	1886(d)(5)(M)" after "1881(b)(13)(A)(ii)".
24	(b) Study and Report on Removing Barriers to
25	Development of New Antimicrobial Drugs.—

1	(1) Study.—The Comptroller General of the
2	United States shall, in consultation with the Director
3	of the National Institutes of Health, the Commissioner
4	of Food and Drugs, and the Director of the Centers
5	for Disease Control and Prevention, conduct a study
6	to—
7	(A) identify and examine the barriers that
8	prevent the development of new antimicrobial
9	drugs, as defined in section $1886(d)(5)(M)(iii)$ of
10	the Social Security Act (42 U.S.C.
11	$1395ww(d)(5)(M)(iii)), \ as \ added \ by \ subsection$
12	(a)(1); and
13	(B) develop recommendations for actions to
14	be taken in order to overcome any barriers iden-
15	tified under subparagraph (A).
16	(2) Report.—Not later than 1 year after the
17	date of the enactment of this Act, the Comptroller
18	General shall submit to Congress a report on the
19	study conducted under paragraph (1).

1	Subtitle H—Vaccine Access,
2	Certainty, and Innovation
3	SEC. 2141. TIMELY REVIEW OF VACCINES BY THE ADVISORY
4	COMMITTEE ON IMMUNIZATION PRACTICES.
5	Section 2102(a) of the Public Health Service Act (42
6	U.S.C. 300aa-2(a)) is amended by adding at the end the
7	following:
8	"(10) Advisory committee on immunization
9	PRACTICES.—
10	"(A) STANDARD PERIODS OF TIME FOR
11	MAKING RECOMMENDATIONS.—Upon the licen-
12	sure of any vaccine or any new indication for a
13	vaccine, the Director of the Program shall direct
14	the Advisory Committee on Immunization Prac-
15	tices, at its next regularly scheduled meeting, to
16	consider the use of the vaccine.
17	"(B) Expedited review pursuant to re-
18	QUEST BY SPONSOR OR MANUFACTURER.—If the
19	Advisory Committee does not make recommenda-
20	tions with respect to the use of a vaccine at the
21	Advisory Committee's first regularly scheduled
22	meeting after the licensure of the vaccine or any
23	new indication for the vaccine, the Advisory
24	Committee, at the request of the sponsor of the

1	vaccine,	shall	make	such	recommendations	on	an
2	expedite	d basi	s.				

"(C) Expedited review for breakThrough therapies and for use during
Public health emergencies.—If a vaccine is
designated as a breakthrough therapy under section 506 of the Federal Food, Drug, and Cosmetic Act and is licensed under section 351 of
this Act, the Advisory Committee shall make recommendations with respect to the use of the vaccine on an expedited basis.

"(D) DEFINITION.—In this paragraph, the terms 'Advisory Committee on Immunization Practices' and 'Advisory Committee' mean the advisory committee on immunization practices established by the Secretary pursuant to section 222, acting through the Director of the Centers for Disease Control and Prevention.".

19 SEC. 2142. REVIEW OF PROCESSES AND CONSISTENCY OF 20 ACIP RECOMMENDATIONS.

21 (a) REVIEW.—The Director of the Centers for Disease 22 Control and Prevention shall conduct a review of the process 23 used by the Advisory Committee on Immunization Practices 24 to evaluate consistency in formulating and issuing rec-25 ommendations pertaining to vaccines.

1	(b) Considerations.—The review under subsection
2	(a) shall include assessment of—
3	(1) the criteria used to evaluate new and existing
4	vaccines;
5	(2) the Grading of Recommendations, Assess-
6	ment, Development, and Evaluation (GRADE) ap-
7	proach to the review and analysis of scientific and
8	economic data, including the scientific basis for such
9	approach; and
10	(3) the extent to which the processes used by the
11	working groups of the Advisory Committee on Immu-
12	nization Practices are consistent among groups.
13	(c) Stakeholders.—In carrying out the review
14	under subsection (a), the Director of the Centers for Disease
15	Control and Prevention shall solicit input from vaccine
16	stakeholders.
17	(d) Report.—Not later than 18 months after the date
18	of enactment of this Act, the Director of the Centers for Dis-
19	ease Control and Prevention shall submit to the appropriate
20	committees of the Congress and make publicly available a
21	report on the results of the review under subsection (a), in-
22	cluding recommendations on improving the consistency of
23	the process described in such subsection.
24	(e) Definition.—In this section, the term "Advisory
25	Committee on Immunization Practices" means the advisory

1	committee on immunization practices established by the
2	Secretary of Health and Human Services pursuant to sec-
3	tion 222 of the Public Health Service Act (42 U.S.C. 217a),
4	acting through the Director of the Centers for Disease Con-
5	trol and Prevention.
6	SEC. 2143. MEETINGS BETWEEN CDC AND VACCINE DEVEL-
7	OPERS.
8	Section 310 of the Public Health Service Act (42
9	U.S.C. 2420) is amended by adding at the end the following:
10	"(c)(1) In this subsection, the term 'vaccine developer'
11	means a nongovernmental entity engaged in—
12	"(A)(i) the development of a vaccine with the in-
13	tent to pursue licensing of the vaccine by the Food
14	and Drug Administration; or
15	"(ii) the production of a vaccine licensed by the
16	Food and Drug Administration; and
17	"(B) vaccine research.
18	"(2)(A) Upon the submission of a written request for
19	a meeting by a vaccine developer, that includes a justifica-
20	tion for the meeting, the Secretary, acting through the Di-
21	rector of the Centers for Disease Control and Prevention,
22	shall convene a meeting of representatives of the vaccine de-
23	veloper and experts from the Centers for Disease Control
24	and Prevention in immunization programs, epidemiology,
25	and other relevant areas at which the Director (or the Di-

- 1 rector's designee), for the purpose of informing the vaccine
- 2 developer's understanding of public health needs and prior-
- 3 ities, shall provide the perspectives of the Centers for Dis-
- 4 ease Control and Prevention and other relevant Federal
- 5 agencies regarding—
- 6 "(i) public health needs, epidemiology, and im-
- 7 plementation considerations with regard to a vaccine
- 8 developer's potential vaccine profile; and
- 9 "(ii) potential implications of such perspectives
- for the vaccine developer's vaccine research and devel-
- 11 opment planning.
- 12 "(B) In addition to the representatives specified in
- 13 subparagraph (A), the Secretary may, with the agreement
- 14 of the vaccine developer requesting a meeting under such
- 15 subparagraph, include in such meeting representatives of—
- 16 "(i) the Food and Drug Administration; and
- "(ii) the National Vaccine Program.
- 18 "(C) The Secretary shall convene a meeting requested
- 19 under subparagraph (A) not later than 120 days after re-
- 20 ceipt of the request for the meeting.
- 21 "(3)(A) Upon the submission of a written request by
- 22 a vaccine developer, the Secretary, acting through the Direc-
- 23 tor of the Centers for Disease Control and Prevention, shall
- 24 provide to the vaccine developer any age-based or other de-

1	mographically assessed disease epidemiological analyses or
2	data that—
3	"(i) are specified in the request;
4	"(ii) have been published;
5	"(iii) have been performed by or are in the pos-
6	session of the Centers;
7	"(iv) are not a trade secret or commercial or fi-
8	nancial information that is privileged or confidential
9	and subject to section 552(b)(4) of title 5, United
10	States Code, or section 1905 of title 18, United States
11	Code; and
12	"(v) do not contain individually identifiable in-
13	formation.
14	"(B) The Secretary shall provide analyses requested by
15	a vaccine manufacturer under subparagraph (A) not later
16	than 120 calendar days after receipt of the request for the
17	analyses.
18	"(4) The Secretary shall promptly notify a vaccine de-
19	veloper if—
20	"(A) the Secretary becomes aware of any change
21	to information that was—
22	"(i) shared by the Secretary with the vac-
23	cine developer during a meeting under para-
24	graph (2); or

1	"(ii) provided by the Secretary to the vac-
2	cine developer in one or more analyses under
3	paragraph (3); and
4	"(B) the change to such information may have
5	implications for the vaccine developer's vaccine re-
6	search and development.".
7	Subtitle I—Orphan Product Exten-
8	sions Now; Incentives for Cer-
9	tain Products for Limited Popu-
10	lations
11	SEC. 2151. EXTENSION OF EXCLUSIVITY PERIODS FOR A
12	DRUG APPROVED FOR A NEW INDICATION
13	FOR A RARE DISEASE OR CONDITION.
14	(a) In General.—Chapter V of the Federal Food,
15	Drug, and Cosmetic Act, as amended by sections 2062 and
16	2063, is further amended by inserting after section 505H
17	of such Act the following:
18	"SEC. 505I. EXTENSION OF EXCLUSIVITY PERIODS FOR A
19	DRUG APPROVED FOR A NEW INDICATION
20	FOR A RARE DISEASE OR CONDITION.
21	"(a) Designation.—
22	"(1) In General.—The Secretary shall des-
23	ignate a drug as a drug approved for a new indica-
24	tion to prevent, diagnose, or treat a rare disease or

1	condition for purposes of granting the extensions
2	under subsection (b) if—
3	"(A) prior to approval of an application or
4	supplemental application for the new indication,
5	the drug was approved or licensed for marketing
6	under section 505(c) of this Act or section 351(a)
7	of the Public Health Service Act, but was not so
8	approved or licensed for the new indication;
9	" $(B)(i)$ the sponsor of the approved or li-
10	censed drug files an application or a supple-
11	mental application for approval of the new indi-
12	cation for use of the drug to prevent, diagnose,
13	or treat the rare disease or condition; and
14	"(ii) the Secretary approves the application
15	or supplemental application; and
16	"(C) the application or supplemental appli-
17	cation for the new indication contains the con-
18	sent of the applicant to notice being given by the
19	Secretary under paragraph (4) respecting the
20	designation of the drug.
21	"(2) Revocation of Designation.—
22	"(A) In general.—Except as provided in
23	subparagraph (B), a designation under para-
24	graph (1) shall not be revoked for any reason.

1	"(B) Exception.—The Secretary may re-
2	voke a designation of a drug under paragraph
3	(1) if the Secretary finds that the application or
4	supplemental application resulting in such des-
5	ignation contained an untrue statement of mate-
6	$rial\ fact.$
7	"(3) Notification prior to discontinuance
8	OF PRODUCTION FOR SOLELY COMMERCIAL REA-
9	SONS.—A designation of a drug under paragraph (1)
10	shall be subject to the condition that the sponsor of the
11	drug will notify the Secretary of any discontinuance
12	of the production of the drug for solely commercial
13	reasons at least one year before such discontinuance.
14	"(4) Notice to public.—Notice respecting the
15	designation of a drug under paragraph (1) shall be
16	made available to the public.
17	"(b) Extension.—If the Secretary designates a drug
18	as a drug approved for a new indication for a rare disease
19	or condition, as described in subsection (a)(1)—
20	" $(1)(A)$ the 4-, 5-, and 7½-year periods de-
21	scribed in subsections $(c)(3)(E)(ii)$ and $(j)(5)(F)(ii)$ of
22	section 505, the 3-year periods described in clauses
23	(iii) and (iv) of subsection $(c)(3)(E)$ and clauses (iii)
24	and (iv) of subsection $(j)(5)(F)$ of section 505, and the

7-year period described in section 527, as applicable,
 shall be extended by 6 months; or

"(B) the 4- and 12-year periods described in subparagraphs (A) and (B) of section 351(k)(7) of the Public Health Service Act and the 7-year period described in section 527, as applicable, shall be extended by 6 months; and

"(2)(A) if the drug is the subject of a listed patent for which a certification has been submitted under subsection (b)(2)(A)(ii) or (j)(2)(A)(vii)(II) of section 505 or a listed patent for which a certification has been submitted under subsections (b)(2)(A)(iii) or (j)(2)(A)(vii)(III) of section 505, the period during which an application may not be approved under section 505(c)(3) or section 505(j)(5)(B) shall be extended by a period of 6 months after the date the patent expires (including any patent extensions); or

"(B) if the drug is the subject of a listed patent for which a certification has been submitted under subsection (b)(2)(A)(iv) or (j)(2)(A)(vii)(IV) of section 505, and in the patent infringement litigation resulting from the certification the court determines that the patent is valid and would be infringed, the period during which an application may not be approved under section 505(c)(3) or section 505(j)(5)(B) shall

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- 1 be extended by a period of 6 months after the date the
- 2 patent expires (including any patent extensions).
- 3 "(c) Relation to Pediatric and Qualified Infec-
- 4 TIOUS DISEASE PRODUCT EXCLUSIVITY.—Any extension
- 5 under subsection (b) of a period shall be in addition to any
- 6 extension of the periods under sections 505A and 505E of
- 7 this Act and section 351(m) of the Public Health Service
- 8 Act, as applicable, with respect to the drug.
- 9 "(d) Limitations.—The extension described in sub-
- 10 section (b) shall not apply if the drug designated under sub-
- 11 section (a)(1) has previously received an extension by oper-
- 12 ation of subsection (b).
- 13 "(e) Definition.—In this section, the term 'rare dis-
- 14 ease or condition' has the meaning given to such term in
- 15 section 526(a)(2).".
- 16 (b) APPLICATION.—Section 505G of the Federal Food,
- 17 Drug, and Cosmetic Act, as added by subsection (a), applies
- 18 only with respect to a drug for which an application or
- 19 supplemental application described in subsection
- 20 (a)(1)(B)(i) of such section 505G is first approved under
- 21 section 505(c) of such Act (21 U.S.C. 355(c)) or section
- 22 351(a) of the Public Health Service Act (42 U.S.C. 262(a))
- 23 on or after the date of the enactment of this Act.
- 24 (c) Conforming Amendments.—

1	(1) Relation to pediatric exclusivity for
2	DRUGS.—Section 505A of the Federal Food, Drug,
3	and Cosmetic Act (21 U.S.C. 355a) is amended—
4	(A) in subsection (b), by adding at the end
5	the following:
6	"(3) Relation to exclusivity for a drug ap-
7	PROVED FOR A NEW INDICATION FOR A RARE DISEASE
8	OR CONDITION.—Notwithstanding the references in
9	paragraph (1) to the lengths of the exclusivity periods
10	after application of pediatric exclusivity, the 6-month
11	extensions described in paragraph (1) shall be in ad-
12	dition to any extensions under section 505G."; and
13	(B) in subsection (c), by adding at the end
14	the following:
15	"(3) Relation to exclusivity for a drug ap-
16	PROVED FOR A NEW INDICATION FOR A RARE DISEASE
17	OR CONDITION.—Notwithstanding the references in
18	paragraph (1) to the lengths of the exclusivity periods
19	after application of pediatric exclusivity, the 6-month
20	extensions described in paragraph (1) shall be in ad-
21	dition to any extensions under section 505G.".
22	(2) Relation to exclusivity for New Quali-
23	FIED INFECTIOUS DISEASE PRODUCTS THAT ARE
24	DRUGS.—Subsection (b) of section 505E of the Fed-

1	eral Food, Drug, and Cosmetic Act (21 U.S.C. 355f)
2	is amended—
3	(A) by amending the subsection heading to
4	read as follows: "Relation to Pediatric Ex-
5	CLUSIVITY AND EXCLUSIVITY FOR A DRUG AP-
6	PROVED FOR A NEW INDICATION FOR A RARE
7	Disease or Condition.—"; and
8	(B) by striking "any extension of the period
9	under section 505A" and inserting "any exten-
10	sion of the periods under sections 505A and
11	505G, as applicable,".
12	(3) Relation to pediatric exclusivity for
13	BIOLOGICAL PRODUCTS.—Section 351(m) of the Pub-
14	lic Health Service Act (42 U.S.C. 262(m)) is amended
15	by adding at the end the following:
16	"(5) Relation to exclusivity for a biologi-
17	CAL PRODUCT APPROVED FOR A NEW INDICATION FOR
18	A RARE DISEASE OR CONDITION.—Notwithstanding
19	the references in paragraphs $(2)(A)$, $(2)(B)$, $(3)(A)$,
20	and (3)(B) to the lengths of the exclusivity periods
21	after application of pediatric exclusivity, the 6-month
22	extensions described in such paragraphs shall be in
23	addition to any extensions under section 505G.".

1	SEC. 2152. REAUTHORIZATION OF RARE PEDIATRIC DIS-
2	EASE PRIORITY REVIEW VOUCHER INCENTIVE
3	PROGRAM.
4	(a) In General.—Section 529 of the Federal Food,
5	Drug, and Cosmetic Act (21 U.S.C. 360ff) is amended—
6	(1) in subsection (a)—
7	(A) in paragraph (3), by amending sub-
8	paragraph (A) to read as follows:
9	"(A) The disease is a serious or life-threat-
10	ening disease in which the serious or life-threat-
11	ening manifestations primarily affect individ-
12	uals aged from birth to 18 years, including age
13	groups often called neonates, infants, children,
14	and adolescents."; and
15	(B) in paragraph (4)—
16	(i) in subparagraph (E), by striking
17	"and" at the end;
18	(ii) in subparagraph (F), by striking
19	the period at the end and inserting "; and";
20	and
21	(iii) by adding at the end the fol-
22	lowing:
23	"(G) is for a drug or biological product for
24	which a priority review voucher has not been
25	issued under section 524 (relating to tropical
26	disease products).": and

1	(2) in subsection (b), by striking paragraph (5)
2	and inserting the following:
3	"(5) Termination of Authority.—The Sec-
4	retary may not award any priority review vouchers
5	under paragraph (1) after December 31, 2018.".
6	(b) GAO STUDY AND REPORT.—
7	(1) Study.—The Comptroller General of the
8	United States shall conduct a study on the effective-
9	ness of awarding priority review vouchers under sec-
10	tion 529 of the Federal Food, Drug, and Cosmetic Act
11	(21 U.S.C. 360ff) in providing incentives for the de-
12	velopment of drugs that treat or prevent rare pedi-
13	atric diseases (as defined in subsection (a)(3) of such
14	section) that would not otherwise have been developed.
15	In conducting such study, the Comptroller General
16	shall examine the following:
17	(A) The indications for which each drug for
18	which a priority review voucher was awarded
19	under such section 529 was approved under sec-
20	tion 505 of such Act (21 U.S.C. 355) or section
21	351 of the Public Health Service Act (42 U.S.C.
22	262).
23	(B) Whether the priority review voucher im-
24	pacted a sponsor's decision to invest in devel-

1	oping a drug to treat or prevent a rare pediatric
2	disease.
3	(C) An analysis of the drugs that utilized
4	such priority review vouchers, which shall in-
5	clude—
6	(i) the indications for which such
7	drugs were approved under section 505 of
8	the Federal Food, Drug, and Cosmetic Act
9	(21 U.S.C. 355) or section 351 of the Public
10	Health Service Act (42 U.S.C. 262);
11	(ii) whether unmet medical needs were
12	addressed through the approval of such
13	drugs, including, for each such drug—
14	(I) if an alternative therapy was
15	previously available to treat the indi-
16	cation; and
17	(II) the benefit or advantage the
18	drug provided over another available
19	the rapy;
20	(iii) the number of patients potentially
21	treated by such drugs;
22	(iv) the value of the priority review
23	voucher if transferred; and

1	(v) the length of time between the date
2	on which a priority review voucher was
3	awarded and the date on which it was used.
4	(D) With respect to the priority review
5	voucher program under section 529 of the Fed-
6	eral Food, Drug, and Cosmetic Act (21 U.S.C.
7	360ff)—
8	(i) the resources used by, and burden
9	placed on, the Food and Drug Administra-
10	tion in implementing such program, includ-
11	ing the effect of such program on the Food
12	and Drug Administration's review of drugs
13	for which a priority review voucher was not
14	awarded or used;
15	(ii) the impact of the program on the
16	public health as a result of the expedited re-
17	view of applications for drugs that treat or
18	prevent non-serious indications that are
19	generally used by the broader public; and
20	(iii) alternative approaches to improv-
21	ing such program so that the program is
22	appropriately targeted toward providing in-
23	centives for the development of clinically
24	important drugs that—

1	(I) prevent or treat rare pediatric
2	diseases; and
3	(II) would likely not otherwise
4	have been developed to prevent or treat
5	such diseases.
6	(2) Report.—Not later than December 31, 2017,
7	the Comptroller General of the United States shall
8	submit to the Committee on Energy and Commerce of
9	the House of Representatives and the Committee on
10	Health, Education, Labor and Pensions of the Senate
11	a report containing the results of the study of con-
12	ducted under paragraph (1).
13	Subtitle J—Domestic Manufac-
14	turing and Export Efficiencies
15	SEC. 2161. GRANTS FOR STUDYING THE PROCESS OF CON-
15 16	
	SEC. 2161. GRANTS FOR STUDYING THE PROCESS OF CON-
16 17	SEC. 2161. GRANTS FOR STUDYING THE PROCESS OF CONTINUOUS DRUG MANUFACTURING.
16 17 18	SEC. 2161. GRANTS FOR STUDYING THE PROCESS OF CONTINUOUS DRUG MANUFACTURING. (a) IN GENERAL.—The Commissioner of Food and
16 17 18 19	SEC. 2161. GRANTS FOR STUDYING THE PROCESS OF CONTINUOUS DRUG MANUFACTURING. (a) IN GENERAL.—The Commissioner of Food and Drugs may award grants to institutions of higher education
16 17 18 19	SEC. 2161. GRANTS FOR STUDYING THE PROCESS OF CON- TINUOUS DRUG MANUFACTURING. (a) In General.—The Commissioner of Food and Drugs may award grants to institutions of higher education and nonprofit organizations for the purpose of studying
16 17 18 19 20	SEC. 2161. GRANTS FOR STUDYING THE PROCESS OF CONTINUOUS DRUG MANUFACTURING. (a) IN GENERAL.—The Commissioner of Food and Drugs may award grants to institutions of higher education and nonprofit organizations for the purpose of studying and recommending improvements to the process of continuous commending improvements to the process of continuous commending improvements.

1	(1) The term "drug" has the meaning given to
2	such term in section 201 of the Federal Food, Drug,
3	and Cosmetic Act (21 U.S.C. 321).
4	(2) The term "biological product" has the mean-
5	ing given to such term in section 351(i) of the Public
6	Health Service Act (42 U.S.C. 262(i)).
7	(3) The term "institution of higher education"
8	has the meaning given to such term in section 101 of
9	the Higher Education Act of 1965 (20 U.S.C. 1001).
10	(c) Authorization of Appropriations.—There is
11	authorized to be appropriated to carry out this section
12	\$5,000,000 for each of fiscal years 2016 through 2020.
13	SEC. 2162. RE-EXPORTATION AMONG MEMBERS OF THE EU-
13 14	SEC. 2162. RE-EXPORTATION AMONG MEMBERS OF THE EU- ROPEAN ECONOMIC AREA.
14	ROPEAN ECONOMIC AREA.
14 15	ROPEAN ECONOMIC AREA. Section 1003 of the Controlled Substances Import and
14 15 16	ROPEAN ECONOMIC AREA. Section 1003 of the Controlled Substances Import and Export Act (21 U.S.C. 953) is amended—
14 15 16 17	ROPEAN ECONOMIC AREA. Section 1003 of the Controlled Substances Import and Export Act (21 U.S.C. 953) is amended— (1) in subsection (f)—
14 15 16 17	ROPEAN ECONOMIC AREA. Section 1003 of the Controlled Substances Import and Export Act (21 U.S.C. 953) is amended— (1) in subsection (f)— (A) in paragraph (5)—
114 115 116 117 118	ROPEAN ECONOMIC AREA. Section 1003 of the Controlled Substances Import and Export Act (21 U.S.C. 953) is amended— (1) in subsection (f)— (A) in paragraph (5)— (i) by striking "(5)" and inserting
14 15 16 17 18 19 20	ROPEAN ECONOMIC AREA. Section 1003 of the Controlled Substances Import and Export Act (21 U.S.C. 953) is amended— (1) in subsection (f)— (A) in paragraph (5)— (i) by striking "(5)" and inserting "(5)(A)";
14 15 16 17 18 19 20 21	ROPEAN ECONOMIC AREA. Section 1003 of the Controlled Substances Import and Export Act (21 U.S.C. 953) is amended— (1) in subsection (f)— (A) in paragraph (5)— (i) by striking "(5)" and inserting "(5)(A)"; (ii) by inserting ", except that the con-
14 15 16 17 18 19 20 21	ROPEAN ECONOMIC AREA. Section 1003 of the Controlled Substances Import and Export Act (21 U.S.C. 953) is amended— (1) in subsection (f)— (A) in paragraph (5)— (i) by striking "(5)" and inserting "(5)(A)"; (ii) by inserting ", except that the controlled substance may be exported from the

1	(iii) by adding at the end the fol-
2	lowing:
3	"(B) Subsequent to any re-exportation described
4	in subparagraph (A), a controlled substance may con-
5	tinue to be exported from any country that is a mem-
6	ber of the European Economic Area to any other such
7	country, provided that—
8	"(i) the conditions applicable with respect
9	to the first country under paragraphs (1), (2),
10	(3), (4), (6), and (7) are met by each subsequent
11	country from which the controlled substance is
12	exported pursuant to this paragraph; and
13	"(ii) the conditions applicable with respect
14	to the second country under such paragraphs are
15	met by each subsequent country to which the con-
16	trolled substance is exported pursuant to this
17	paragraph."; and
18	(B) in paragraph (6)—
19	(i) by striking "(6)" and inserting
20	"(6)(A)"; and
21	(ii) by adding at the end the following:
22	"(B) In the case of re-exportation among mem-
23	bers of the European Economic Area, within 30 days
24	after each re-exportation, the person who exported the

1	controlled substance from the United States delivers to
2	the Attorney General—
3	"(i) documentation certifying that such re-
4	exportation has occurred; and
5	"(ii) information concerning the consignee,
6	country, and product."; and
7	(2) by adding at the end the following:
8	"(g) Limitation.—The Attorney General shall not
9	promulgate nor enforce any regulation, subregulatory guid-
10	ance, or enforcement policy which impedes re-exportation
11	among European Economic Area countries (as provided in
12	subsection (f)(5)), including by promulgating or enforcing
13	any requirement that—
14	"(1) re-exportation from the first country to the
15	second country or re-exportation from the second
16	country to another country (as such terms are used
17	in subsection (f)) occur within a specified period of
18	$time;\ or$
19	"(2) information concerning the consignee, coun-
20	try, and product be provided prior to exportation of
21	the controlled substance from the United States or
22	prior to each re-exportation among members of the
23	European Economic Area.".

1	Subtitle K—Enhancing
2	Combination Products Review
3	SEC. 2181. ENHANCING COMBINATION PRODUCTS REVIEW.
4	Section $503(g)(4)(C)$ of the Federal Food, Drug, and
5	Cosmetic Act (21 U.S.C. 353(g)(4)(C)) is amended by add-
6	ing at the end the following new clause:
7	"(iii) Not later than 18 months after the date of the
8	enactment of the 21st Century Cures Act, the Secretary shall
9	issue final guidance that describes the responsibilities of
10	each agency center regarding its review of combination
11	products. The Secretary shall, after soliciting public com-
12	ment, review and update the guidance periodically.".
13	Subtitle L—Priority Review for
14	Breakthrough Devices
15	SEC. 2201. PRIORITY REVIEW FOR BREAKTHROUGH DE-
16	VICES.
17	(a) In General.—Chapter V of the Federal Food,
18	Drug, and Cosmetic Act is amended—
19	(1) in section $515(d)$ —
20	(A) by striking paragraph (5); and
21	(B) by redesignating paragraph (6) as
22	paragraph (5); and
23	(2) by inserting after section 515A (21 U.S.C.
24	360e-1) the following:

1	"SEC. 515B. PRIORITY REVIEW FOR BREAKTHROUGH DE-
2	VICES.
3	"(a) In General.—In order to provide for more effec-
4	tive treatment or diagnosis of life-threatening or irrevers-
5	ibly debilitating human diseases or conditions, the Sec-
6	retary shall establish a program to provide priority review
7	for devices—
8	"(1) representing breakthrough technologies;
9	"(2) for which no approved alternatives exist;
10	"(3) offering significant advantages over existing
11	approved or cleared alternatives, including the poten-
12	tial to, compared to existing approved or cleared al-
13	ternatives, reduce or eliminate the need for hos-
14	pitalization, improve patient quality of life, facilitate
15	patients' ability to manage their own care (such as
16	through self-directed personal assistance), or establish
17	long-term clinical efficiencies; or
18	"(4) the availability of which is in the best inter-
19	est of patients.
20	"(b) Request for Designation.—A sponsor of a de-
21	vice may request that the Secretary designate the device for
22	priority review under this section. Any such request for des-
23	ignation may be made at any time prior to the submission
24	of an application under section 515(c), a petition for classi-
25	fication under section $513(f)(2)$, or a notification under sec-
26	tion 510(k).

1	"(c) Designation Process.—
2	"(1) In general.—Not later than 60 calendar
3	days after the receipt of a request under subsection
4	(b), the Secretary shall determine whether the device
5	that is the subject of the request meets the criteria de-
6	scribed in subsection (a). If the Secretary determines
7	that the device meets the criteria, the Secretary shall
8	designate the device for priority review.
9	"(2) REVIEW.—Review of a request under sub-
10	section (b) shall be undertaken by a team that is com-
11	posed of experienced staff and managers of the Food
12	and Drug Administration and is chaired by a senior
13	manager.
14	"(3) Designation determination.—A deter-
15	mination approving or denying a request under sub-
16	section (b) shall be considered a significant decision
17	under section 517A and the Secretary shall provide a
18	written, substantive summary of the basis for the de-
19	termination in accordance with section $517A(a)$.
20	"(4) Reconsideration.—
21	"(A) REQUEST FOR RECONSIDERATION.—
22	Any person whose request under subsection (b) is
23	denied may, within 30 days of the denial, re-
24	quest reconsideration of the denial in accordance

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with section 517A(b)—

1	"(i) based upon the submission of doc-
2	uments by such person; or
3	"(ii) based upon such documents and a
4	meeting or teleconference.
5	"(B) Response.—Reconsideration of a des-
6	ignation determination under this paragraph
7	shall be conducted in accordance with section
8	517A(b).
9	"(5) Withdrawal.—If the Secretary approves a
10	priority review designation for a device under this
11	section, the Secretary may not withdraw the designa-
12	tion based on the fact that the criteria specified in
13	subsection (a) are no longer met because of the subse-
14	quent clearance or approval of another device that
15	was designated under—
16	"(A) this section; or
17	"(B) section $515(d)(5)$ (as in effect imme-
18	diately prior to the enactment of the 21st Cen-
19	tury Cures Act).
20	"(d) Priority Review.—
21	"(1) Actions.—For purposes of expediting the
22	development and review of devices designated under
23	subsection (c), the Secretary shall—
24	"(A) assign a team of staff, including a
25	team leader with appropriate subject matter ex-

1	pertise and experience, for each device for which
2	a request is submitted under subsection (b);
3	"(B) provide for oversight of the team by
4	senior agency personnel to facilitate the efficient
5	development of the device and the efficient review
6	of any submission described in subsection (b) for
7	the device;
8	"(C) adopt an efficient process for timely
9	$dispute \ resolution;$
10	"(D) provide for interactive communication
11	with the sponsor of the device during the review
12	process;
13	"(E) expedite the Secretary's review of
14	manufacturing and quality systems compliance,
15	as applicable;
16	"(F) disclose to the sponsor in advance the
17	topics of any consultation concerning the spon-
18	sor's device that the Secretary intends to under-
19	take with external experts or an advisory com-
20	mittee and provide the sponsor an opportunity
21	to recommend such external experts;
22	"(G) for applications submitted under sec-
23	tion 515(c), provide for advisory committee
24	input, as the Secretary determines appropriate

1	(including in response to the request of the spon-
2	sor); and
3	"(H) assign staff to be available within a
4	reasonable time to address questions posed by in-
5	stitutional review committees concerning the con-
6	ditions and clinical testing requirements appli-
7	cable to the investigational use of the device pur-
8	suant to an exemption under section $520(g)$.
9	"(2) Additional Actions.—In addition to the
10	actions described in paragraph (1), for purposes of
11	expediting the development and review of devices des-
12	ignated under subsection (c), the Secretary, in col-
13	laboration with the device sponsor, may, as appro-
14	priate—
15	"(A) coordinate with the sponsor regarding
16	early agreement on a data development plan;
17	"(B) take steps to ensure that the design of
18	clinical trials is as efficient as practicable, such
19	as through adoption of shorter or smaller clinical
20	trials, application of surrogate endpoints, and
21	use of adaptive trial designs and Bayesian sta-
22	tistics, to the extent scientifically appropriate;
23	"(C) facilitate, to the extent scientifically
24	appropriate, expedited and efficient development
25	and review of the device through utilization of

1	timely postmarket data collection, with regard to
2	applications for approval under section $515(c)$;
3	and
4	"(D) agree to clinical protocols that the Sec-
5	retary will consider binding on the Secretary
6	and the sponsor, subject to—
7	"(i) changes agreed to by the sponsor
8	and the Secretary;
9	"(ii) changes that the Secretary deter-
10	mines are required to prevent an unreason-
11	able risk to the public health; or
12	"(iii) the identification of a substan-
13	tial scientific issue determined by the Sec-
14	retary to be essential to the safety or effec-
15	tiveness of the device involved.
16	"(e) Priority Review Guidance.—
17	"(1) Content.—The Secretary shall issue guid-
18	ance on the implementation of this section. Such
19	guidance shall include the following:
20	"(A) The process for a person to seek a pri-
21	ority review designation.
22	"(B) A template for requests under sub-
23	section (b).
24	"(C) The criteria the Secretary will use in
25	evaluating a request for priority review.

1 "(D) The standards the Secretary will use 2 in assigning a team of staff, including team 3 leaders, to review devices designated for priority 4 review, including any training required for such 5 personnel on effective and efficient review.

> "(2) Process.—Prior to finalizing the guidance under paragraph (1), the Secretary shall propose such guidance for public comment.

"(f) Construction.—

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- "(1) Purpose.—This section is intended to encourage the Secretary and provide the Secretary sufficient authorities to apply efficient and flexible approaches to expedite the development of, and prioritize the agency's review of, devices that represent breakthrough technologies.
- "(2) Construction.—Nothing in this section shall be construed to alter the criteria and standards for evaluating an application pursuant to section 515(c), a report and request for classification under section 513(f)(2), or a report under section 510(k), including the recognition of valid scientific evidence as described in section 513(a)(3)(B), and consideration of the least burdensome means of evaluating device effectiveness or demonstrating substantial equivalence between devices with differing technological character-

1	istics, as applicable. Nothing in this section alters the
2	authority of the Secretary to act on an application
3	pursuant to section 515(d) before completion of an es-
4	tablishment inspection, as the Secretary deems appro-
5	priate.".
6	(b) Conforming Amendment Related to Designa-
7	TION DETERMINATIONS.—Section 517A(a)(1) of the Federal
8	Food, Drug, and Cosmetic Act (21 U.S.C. 360g-1(a)(1)) is
9	amended by inserting "a request for designation under sec-
10	tion 515B," after "an application under section 515,".
11	Subtitle M—Medical Device
12	Regulatory Process Improvements
13	SEC. 2221. THIRD-PARTY QUALITY SYSTEM ASSESSMENT.
14	(a) Establishment of Third-Party Quality Sys-
15	TEM ASSESSMENT PROGRAM.—Chapter V of the Federal
16	Food, Drug, and Cosmetic Act is amended by inserting after
17	section 524A (21 U.S.C. 360n-1) the following new section.
18	"SEC. 524B. THIRD-PARTY QUALITY SYSTEM ASSESSMENT.
19	"(a) Accreditation and Assessment.—
20	"(1) In general; certification of device
21	QUALITY SYSTEM.—The Secretary shall, in accordance
22	with this section, establish a third-party quality sys-
23	tem assessment program—
24	"(A) to accredit persons to assess whether a
25	requestor's quality system, including its design

1	controls, can reasonably assure the safety and ef-
2	fectiveness of in-scope devices subject to device-re-
3	lated changes;
4	"(B) under which accredited persons shall
5	(as applicable) certify that a requestor's quality
6	system meets the criteria included in the guid-
7	ance issued under paragraph (5) with respect to
8	the in-scope devices at issue; and
9	"(C) under which the Secretary shall rely
10	on such certifications for purposes of deter-
11	mining the safety and effectiveness (or as appli-
12	cable, substantial equivalence) of in-scope devices
13	subject to the device-related changes involved, in
14	lieu of compliance with the following submission
15	requirements:
16	"(i) A premarket notification.
17	$``(ii)\ A\ thirty-day\ notice.$
18	"(iii) A Special PMA supplement.
19	"(2) Definitions.—For purposes of this sec-
20	tion-
21	"(A) the term 'device-related changes' means
22	changes made by a requestor with respect to in-
23	scope devices, which are—
24	"(i) changes to a device found to be
25	substantially equivalent under sections

1	513(i) and 510(k) to a predicate device,
2	that—
3	"(I) would otherwise be subject to
4	a premarket notification; and
5	"(II) do not alter—
6	"(aa) the intended use of the
7	changed device; or
8	"(bb) the fundamental sci-
9	entific technology of such device;
10	"(ii) manufacturing changes subject to
11	a 30-day notice;
12	"(iii) changes that qualify for a Spe-
13	cial PMA Supplement; and
14	"(iv) such other changes relating to the
15	devices or the device manufacturing process
16	as the Secretary determines appropriate;
17	"(B) the term 'in-scope device' means a de-
18	vice within the scope of devices agreed to by the
19	requestor and the accredited person for purposes
20	of a request for certification under this section;
21	"(C) the term 'premarket notification'
22	means a premarket notification under section
23	510(k);
24	"(D) the term 'quality system' means the
25	methods used in, and the facilities and controls

1	used for, the design, manufacture, packaging, la-
2	beling, storage, installation, and servicing of de-
3	vices, as described in section 520(f);
4	"(E) the term 'requestor' means a device
5	manufacturer that is seeking certification under
6	this section of a quality system used by such
7	manufacturer;
8	"(F) the term 'Special PMA' means a Spe-
9	cial PMA supplement under section 814.39(d) of
10	title 21, Code of Federal Regulations (or any
11	successor regulations); and
12	"(G) the term 'thirty-day notice' means a
13	notice described in section $515(d)(6)$.
14	"(3) Accreditation process; accreditation
15	RENEWAL.—Except as inconsistent with this section,
16	the process and qualifications for accreditation of per-
17	sons and renewal of such accreditation under section
18	704(g) shall apply with respect to accreditation of
19	persons and renewal of such accreditation under this
20	section.
21	"(4) Use of accredited parties to conduct
22	ASSESSMENTS.—
23	"(A) Initiation of assessment serv-
24	ICES.—

1	"(i) Date assessments author-
2	IZED.—Beginning after the date on which
3	the final guidance is issued under para-
4	graph (5), an accredited person may con-
5	duct an assessment under this section.
6	"(ii) Initiation of assessments.—
7	Use of one or more accredited persons to as-
8	sess a requestor's quality system under this
9	section with respect to in-scope devices shall
10	be at the initiation of the person who reg-
11	isters and lists the devices at issue under
12	section 510.
13	"(B) Compensation for
14	such accredited persons shall—
15	"(i) be determined by agreement be-
16	tween the accredited person and the person
17	who engages the services of the accredited
18	person; and
19	"(ii) be paid by the person who en-
20	gages such services.
21	"(C) Accredited person selection.—
22	Each person who chooses to use an accredited
23	person to assess a requestor's quality system, as
24	described in this section, shall select the accred-
25	ited person from a list of such persons published

1	by the Secretary in accordance with section
2	704(g)(4).
3	"(5) Guidance; criteria for certifi-
4	CATION.—
5	"(A) In general.—The criteria for certifi-
6	cation of a quality system under this section
7	shall be as specified by the Secretary in guidance
8	issued under this paragraph.
9	"(B) Contents; Certification Cri-
10	TERIA.—The guidance under this paragraph
11	shall include specification of—
12	"(i) evaluative criteria to be used by
13	an accredited person to assess and, as ap-
14	plicable, certify a requestor's quality system
15	under this section with respect to in-scope
16	devices; and
17	"(ii) criteria for accredited persons to
18	apply for a waiver of, and exemptions from,
19	the certification criteria under clause (i).
20	"(C) Timeframe for issuing guid-
21	ANCE.—The Secretary shall issue under this
22	paragraph—
23	"(i) draft guidance not later than 12
24	months after the enactment of the 21st Cen-
25	tury Cures Act; and

1	"(ii) final guidance not later than 12
2	months after issuance of the draft guidance
3	under clause (i).
4	"(b) Use of Third-Party Assessment.—
5	"(1) Assessment summary; certification.—
6	"(A) Submission of assessment to sec-
7	RETARY.—An accredited person who assesses a
8	requestor's quality system under subsection (a)
9	shall submit to the Secretary a summary of the
10	assessment—
11	"(i) within 30 days of the assessment;
12	and
13	"(ii) which shall include (as applica-
14	<i>ble)</i> —
15	"(I) the accredited person's certifi-
16	cation that the requestor has satisfied
17	the criteria specified in the guidance
18	issued under subsection $(a)(5)$ for qual-
19	ity system certification with respect to
20	the in-scope devices at issue; and
21	"(II) any waivers or exemptions
22	from such criteria applied by the ac-
23	credited person.
24	"(B) Treatment of assessments.—Sub-
25	ject to action by the Secretary under subpara-

1	graph (C), with respect to assessments which in-
2	clude a certification under this section—
3	"(i) the Secretary's review of the as-
4	sessment summary shall be deemed complete
5	on the day that is 30 days after the date on
6	which the Secretary receives the summary
7	under subparagraph (A); and
8	"(ii) the assessment summary and cer-
9	tification of the quality system of a re-
10	questor shall be deemed accepted by the Sec-
11	retary on such 30th day.
12	"(C) Actions by Secretary.—
13	"(i) In general.—Within 30 days of
14	receiving an assessment summary and cer-
15	tification under subparagraph (A), the Sec-
16	retary may, by written notice to the accred-
17	ited person submitting such assessment cer-
18	tification, deem any such certification to be
19	provisional beyond such 30-day period, sus-
20	pended pending further review by the Sec-
21	retary, or otherwise qualified or cancelled,
22	based on the Secretary's determination that
23	(as applicable)—
24	``(I) additional information is
25	needed to support such certification;

1	"(II) such assessment or certifi-
2	cation is unwarranted; or
3	"(III) such action with regard to
4	the certification is otherwise justified
5	according to such factors and criteria
6	as the Secretary finds appropriate.
7	"(ii) Acceptance of certifi-
8	CATION.—If following action by the Sec-
9	retary under clause (i) with respect to a
10	certification, the Secretary determines that
11	such certification is acceptable, the Sec-
12	retary shall issue written notice to the ap-
13	plicable accredited person indicating such
14	acceptance.
15	"(2) Notifications to secretary by cer-
16	TIFIED REQUESTORS OR ACCREDITED PERSONS FOR
17	PROGRAM EVALUATION PURPOSES.—
18	"(A) Annual summary report for de-
19	VICE-RELATED CHANGES OTHERWISE SUBJECT
20	TO PREMARKET NOTIFICATION.—A requestor
21	whose quality system is certified under this sec-
22	tion that effectuates device-related changes with
23	respect to in-scope devices, without prior submis-
24	sion of a premarket notification, shall ensure

1	that an annual summary report is submitted to
2	the Secretary by the accredited person which—
3	"(i) describes the changes made to the
4	in-scope device; and
5	"(ii) indicates the effective dates of
6	such changes.
7	"(B) Periodic notification for manu-
8	FACTURING CHANGES OTHERWISE SUBJECT TO
9	THIRTY-DAY NOTICE.—A requestor whose quality
10	system is certified under this section that effec-
11	tuates device-related changes with respect to in-
12	scope devices, without prior submission of a thir-
13	ty-day notice, shall provide notification to the
14	Secretary of such changes in the requestor's next
15	periodic report under section 814.84(b) of title
16	21, Code of Federal Regulations (or any suc-
17	cessor regulation). Such notification shall—
18	"(i) describe the changes made; and
19	"(ii) indicate the effective dates of such
20	changes.
21	"(C) Periodic notification for device-
22	RELATED CHANGES OTHERWISE SUBJECT TO
23	SPECIAL PMA SUPPLEMENT.—A requestor whose
24	quality system is certified under this section that
25	effectuates device-related changes with respect to

1	in-scope devices, without prior submission of a
2	Special PMA Supplement, shall provide notifica-
3	tion to the Secretary of such changes in the re-
4	questor's next periodic report under section
5	814.84(b) of title 21, Code of Federal Regulations
6	(or any successor regulation). Such notification
7	shall—
8	"(i) describe the changes made, includ-
9	ing a full explanation of the basis for the
10	changes; and
11	"(ii) indicate the effective dates of such
12	changes.
13	"(D) Use of notifications for program
14	${\it EVALUATION~PURPOSESInformation~submitted}$
15	to the Secretary under subparagraphs (A)
16	through (C) shall be used by the Secretary for
17	purposes of the program evaluation under sub-
18	section (d).
19	"(c) Duration and Effect of Certification.—A
20	certification under this section—
21	"(1) shall remain in effect for a period of 2 years
22	from the date such certification is accepted by the
23	Secretary, subject to paragraph (6);
24	"(2) may be renewed through the process de-
25	scribed in subsection $(a)(3)$;

1	"(3) shall continue to apply with respect to de-
2	vice-related changes made during such 2-year period,
3	provided the certification remains in effect, irrespec-
4	tive of whether such certification is renewed after such
5	2-year period;
6	"(4) shall have no effect on the need to comply
7	with applicable submission requirements specified in
8	subsection (a)(1)(C) with respect to any change per-
9	taining to in-scope devices which is not a device-re-
10	lated change under subsection $(a)(2)$;
11	"(5) shall have no effect on the authority of the
12	Secretary to conduct an inspection or otherwise deter-
13	mine whether the requestor has complied with the ap-
14	plicable requirements of this Act; and
15	"(6) may be revoked by the Secretary upon a de-
16	termination that the requestor's quality system no
17	longer meets the certification criteria specified in the
18	guidance issued under subsection (a)(5) with respect

20 "(d) Notice of Revocation.—The Secretary shall

to the in-scope devices at issue.

 $21 \quad provide \ written \ notification \ to \ the \ requestor \ of \ a \ revocation$

22 pursuant to subsection (c)(6) not later than 10 business

23 days after the determination described in such subsection.

24 Upon receipt of the written notification, the requestor shall

25 satisfy the applicable submission requirements specified in

19

1	subsection (a)(1)(C) for any device-related changes effec-
2	tuated after the date of such determination. After such rev-
3	ocation, such requestor is eligible to seek re-certification
4	under this section of its quality system.
5	"(e) Program Evaluation; Sunset.—
6	"(1) Program evaluation and report.—
7	"(A) EVALUATION.—The Secretary shall
8	complete an evaluation of the third-party quality
9	system assessment program under this section no
10	later than January 31, 2021, based on—
11	"(i) analysis of information from a
12	representative group of device manufactur-
13	ers obtained from notifications provided by
14	certified requestors or accredited persons
15	under subsection (b)(2); and
16	"(ii) such other available information
17	and data as the Secretary determines ap-
18	propriate.
19	"(B) Report.—No later than 1 year after
20	completing the evaluation under subparagraph
21	(A), the Secretary shall issue a report of the eval-
22	uation's findings on the website of the Food and
23	Drug Administration, which shall include the
24	Secretary's recommendations with respect to con-

1	tinuation and as applicable expansion of the
2	program under this section to encompass—
3	"(i) device submissions beyond those
4	$identified\ in\ subsection\ (a)(1)(C);\ and$
5	"(ii) device changes beyond those de-
6	scribed in subsection $(a)(2)(A)$.
7	"(2) Sunset.—This section shall cease to be ef-
8	fective October 1, 2022.
9	"(f) Rule of Construction.—Nothing in this sec-
10	tion shall be construed to limit the authority of the Sec-
11	retary to request and review the complete assessment of a
12	certified requestor under this section on a for-cause basis.".
13	(b) Conforming Amendments.—
14	(1) Requirements for premarket approval
15	SUPPLEMENTS.—Section $515(d)(5)(A)(i)$ of the Fed-
16	eral Food, Drug, and Cosmetic Act (21 U.S.C.
17	360e(d)(5)(A)(i)), as redesignated by section 2201, is
18	further amended by inserting ", subject to section
19	524B" after "that affects safety or effectiveness".
20	(2) Requirements for thirty-day notice.—
21	Section $515(d)(5)(A)(ii)$ of the Federal Food, Drug,
22	and Cosmetic Act (21 U.S.C. $360e(d)(5)(A)(ii)$), as
23	redesignated by section 2201, is further amended by
24	inserting ", subject to section 524B" after "the date
25	on which the Secretary receives the notice".

1	(3) Requirements for premarket notifica-
2	TION; TECHNICAL CORRECTION TO REFERENCE TO
3	SECTION 510(K).—Section 510(l) of the Federal Food,
4	Drug, and Cosmetic Act (21 U.S.C. 360(l)) is amend-
5	ed by striking "of this subsection under subsection
6	(m)" and inserting "of subsection (k) under sub-
7	section (m) or section 524B".
8	(4) Misbranded Devices.—Section 502(t) of
9	the Federal Food, Drug, and Cosmetic Act (21 U.S.C.
10	352(t)) is amended by inserting "or 524B" after "sec-
11	tion 519".
12	SEC. 2222. VALID SCIENTIFIC EVIDENCE.
13	Section 513(a)(3)(B) of the Federal Food, Drug, and
14	Cosmetic Act (21 U.S.C. 360c(a)(3)(B)) is amended—
15	(1) by redesignating clauses (i) and (ii) as sub-
16	clauses (I) and (II), respectively;
17	(2) by striking "(B) If the Secretary" and insert-
18	ing "(B)(i) If the Secretary"; and
19	(3) by adding at the end the following:
20	"(ii) For purposes of clause (i), valid scientific evi-
21	dence may include—
22	"(I) evidence described in well-documented case
23	histories, including registry data, that are collected
24	and monitored under an acceptable protocol;

1	"(II) studies published in peer-reviewed journals;
2	and
3	"(III) data collected in countries other than the
4	United States so long as such data otherwise meet the
5	criteria specified in this subparagraph.
6	"(iii) In the case of a study published in a peer-re-
7	viewed journal that is offered as valid scientific evidence
8	for purposes of clause (i), the Secretary may request data
9	underlying the study if—
10	"(I) the Secretary, in making such request, com-
11	plies with the requirement of subparagraph $(D)(ii)$ to
12	consider the least burdensome appropriate means of
13	$evaluating \ device \ effectiveness \ or \ subsection \ (i)(1)(D)$
14	to consider the least burdensome means of deter-
15	mining substantial equivalence, as applicable;
16	"(II) the Secretary furnishes a written rationale
17	for so requesting the underlying data together with
18	such request; and
19	"(III) if the requested underlying data for such
20	a study are unavailable, the Secretary shall consider
21	such study to be part of the totality of the evidence
22	with respect to the device, as the Secretary determines
23	appropriate.".

1	SEC. 2223. TRAINING AND OVERSIGHT IN LEAST BURDEN-
2	SOME APPROPRIATE MEANS CONCEPT.
3	(a) In General.—Section 513 of the Federal Food,
4	Drug, and Cosmetic Act (21 U.S.C. 360c) is amended by
5	adding at the end the following:
6	"(j) Training and Oversight in Least Burden-
7	Some Appropriate Means Concept.—
8	"(1) Training.—Each employee of the Food and
9	Drug Administration who is involved in the review of
10	premarket submissions under section 515 or section
11	510(k), including supervisors, shall receive training
12	regarding the meaning and implementation of the
13	least burdensome appropriate means concept in the
14	context of the use of that term in subsections
15	(a)(3)(D) and $(i)(1)(D)$ of this section and in section
16	515(c)(5).
17	"(2) Guidance documents.—
18	"(A) Draft updated guidance.—Not
19	later than 12 months after the date of enactment
20	of the 21st Century Cures Act, the Secretary
21	shall issue a draft guidance document updating
22	the October 4, 2002, guidance document entitled
23	'The Least Burdensome Provision of the FDA
24	Modernization Act of 1997: Concept and Prin-
25	ciples; Final Guidance for FDA and Industry'.

1	"(B) Meeting of Stakeholders.—In de-
2	veloping such draft guidance document, the Sec-
3	retary shall convene a meeting of stakeholders to
4	ensure a full record to support the publication of
5	$such\ document.$
6	"(3) Ombudsman Audit.—Not later than 18
7	months after the date of issuance of final version of
8	the draft guidance under paragraph (2), the ombuds-
9	man for the organizational unit of the Food and
10	Drug Administration responsible for the premarket
11	review of devices shall—
12	"(A) conduct, or have conducted, an audit
13	of the training described in paragraph (1); and
14	"(B) include in such audit interviews with
15	a representative sample of persons from industry
16	regarding their experience in the device pre-
17	market review process.".
18	(b) Additional Information Regarding Pre-
19	MARKET APPLICATIONS.—Subsection (c) of section 515 of
20	the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 360e)
21	is amended by adding at the end the following:
22	"(5)(A) Whenever the Secretary requests additional in-
23	formation from an applicant regarding an application
24	under paragraph (1), the Secretary shall consider the least
25	burdensome appropriate means necessary to demonstrate

1	device safety and effectiveness, and request information ac-
2	cordingly.
3	"(B) For purposes of subparagraph (A), the term 'nec-
4	essary' means the minimum required information that
5	would support a determination by the Secretary that an
6	application provides a reasonable assurance of the safety
7	and effectiveness of the device.
8	"(C) Nothing in this paragraph alters the standards
9	for premarket approval of a device.".
10	SEC. 2224. RECOGNITION OF STANDARDS.
11	Section 514(c) of the Federal Food, Drug, and Cos-
12	metic Act (21 U.S.C. 360d(c)) is amended—
13	(1) in paragraph (1), by inserting after subpara-
14	graph (B) the following new subparagraphs:
15	"(C)(i) Any person may submit a request for recogni-
16	tion under subparagraph (A) of all or part of an appro-
17	priate standard established by a nationally or internation-
18	ally recognized standard organization.
19	"(ii) Not later than 60 days after the Secretary re-
20	ceives such a request, the Secretary shall—
21	"(I) make a determination to recognize all, part,
22	or none of the standard that is the subject of the re-
23	quest; and
24	"(II) issue to the person who submitted such re-
25	quest a response in writing that states the Secretary's

1	rationale for that determination, including the sci-
2	entific, technical, regulatory, or other basis for such
3	determination.
4	"(iii) The Secretary shall make a response issued
5	under clause (ii)(II) publicly available, in such manner as
6	the Secretary determines appropriate.
7	"(iv) The Secretary shall take such actions as may be
8	necessary to implement all or part of a standard recognized
9	under clause (i)(I), in accordance with subparagraph (A).
10	"(D) The Secretary shall make publicly available, in
11	such manner as the Secretary determines appropriate, the
12	rationale for recognition under subparagraph (A) of part
13	of a standard, including the scientific, technical, regulatory,
14	or other basis for such recognition."; and
15	(2) by adding at the end the following new para-
16	graphs:
17	"(4) Training on use of standards.—The
18	Secretary shall provide to all employees of the Food
19	and Drug Administration who review premarket sub-
20	missions for devices periodic training on the concept
21	and use of recognized standards for purposes of meet-
22	ing a premarket submission requirement or other ap-
23	plicable requirement under this Act, including stand-
24	ards relevant to an employee's area of device review.
25	"(5) GUIDANCE.—

2 shall publ	ish guidance identifying the principles
3 for recogn	izing standards under this section. In
4 publishing	such guidance, the Secretary shall
5 consider—	-
6	"(i) the experience with, and reliance
7 on, a	standard by other Federal regulatory
8 autho	orities and the device industry; and
9	"(ii) whether recognition of a standard
10 will	promote harmonization among regu-
11 latory	y authorities in the regulation of de-
12 vices.	
13 "(B)	Timing.—The Secretary shall pub-
14 lish—	
15	"(i) draft guidance under subpara-
16 graph	h (A) not later than 12 months after
the d	late of the enactment of the 21st Cen-
18 tury	Cures Act; and
19	"(ii) final guidance not later than 12
20 mont	hs after the close of the public comment
21 perio	d for the draft guidance under clause
22 (i).".	

1	SEC. 2225. EASING REGULATORY BURDEN WITH RESPECT
2	TO CERTAIN CLASS I AND CLASS II DEVICES.
3	(a) Class I Devices.—Section 510(l) of the Federal
4	Food, Drug, and Cosmetic Act (21 U.S.C. 360(l)) is amend-
5	ed—
6	(1) by striking "A report under subsection (k)"
7	and inserting "(1) A report under subsection (k)";
8	and
9	(2) by adding at the end the following new para-
10	graph:
11	"(2) Not later than 120 days after the date of the en-
12	actment of the 21st Century Cures Act, the Secretary shall
13	identify, through publication in the Federal Register, any
14	type of class I device that the Secretary determines no
15	longer requires a report under subsection (k) to provide rea-
16	sonable assurance of safety and effectiveness. Upon such
17	publication—
18	"(A) each type of class I device so identified shall
19	be exempt from the requirement for a report under
20	subsection (k); and
21	"(B) the classification regulation applicable to
22	each such type of device shall be deemed amended to
23	incorporate such exemption.".
24	(b) Class II Devices.—Section 510(m) of the Federal
25	Food, Drug, and Cosmetic Act (21 U.S.C. 360(m)) is
26	amended—

1	(1) by striking paragraph (1) and inserting the
2	following new paragraph: "(1) The Secretary shall—
3	"(A) not later than 60 days after the date of the
4	enactment of the 21st Century Cures Act—
5	"(i) publish in the Federal Register a notice
6	that contains a list of each type of class II device
7	that the Secretary determines no longer requires
8	a report under subsection (k) to provide reason-
9	able assurance of safety and effectiveness; and
10	"(ii) provide for a period of not less than
11	60 days for public comment beginning on the
12	date of the publication of such notice; and
13	"(B) not later than 180 days after the date of the
14	enactment of 21st Century Cures Act, publish in the
15	Federal Register a list representing the Secretary's
16	final determination with respect to the devices in-
17	cluded in the list published under subparagraph
18	(A).";
19	(2) in paragraph (2)—
20	(A) by striking "1 day after the date of the
21	publication of a list under this subsection," and
22	inserting "1 day after the date of publication of
23	the final list under paragraph (1)(B),"; and
24	(B) by striking "30-day period" and insert-
25	ing "60-day period"; and

1	(3) by adding at the end the following new para-
2	graph:
3	"(3) Upon the publication of the final list under para-
4	graph (1)(B)—
5	"(A) each type of class II device so listed shall
6	be exempt from the requirement for a report under
7	subsection (k); and
8	"(B) the classification regulation applicable to
9	each such type of device shall be deemed amended to
10	incorporate such exemption.".
11	SEC. 2226. ADVISORY COMMITTEE PROCESS.
12	(a) Classification Panels.—Paragraph (5) of sec-
13	tion 513(b) of the Federal Food, Drug, and Cosmetic Act
14	(21 U.S.C. 360c(b)) is amended—
15	(1) by striking "(5)" and inserting "(5)(A)";
16	and
17	(2) by adding at the end the following:
18	"(B) When a device is specifically the subject of review
19	by a classification panel, the Secretary shall—
20	"(i) ensure that adequate expertise is represented
21	on the classification panel to assess—
22	"(I) the disease or condition which the de-
23	vice is intended to cure, treat, mitigate, prevent,
24	or diagnose; and
25	"(II) the technology of the device; and

1	"(ii) as part of the process to ensure adequate ex-
2	pertise under clause (i), give due consideration to the
3	recommendations of the person whose premarket sub-
4	mission is subject to panel review on the expertise
5	needed among the voting members of the panel.
6	"(C) For review by a classification panel of a pre-
7	market submission for a device, the Secretary shall—
8	"(i) provide an opportunity for the person whose
9	premarket submission is subject to panel review to
10	provide recommendations on the expertise needed
11	among the voting members of the panel; and
12	"(ii) give due consideration to such recommenda-
13	tions and ensure that adequate expertise is rep-
14	resented on advisory panels to assess—
15	"(I) the disease or condition for which the
16	device is intended to cure, treat, mitigate, pre-
17	vent, or diagnose; and
18	"(II) the technology of the device.
19	"(D) For purposes of subparagraph (B)(ii), the term
20	'adequate expertise' means, with respect to the membership
21	of the classification panel reviewing a premarket submis-
22	sion, that such membership includes—
23	"(i) two or more voting members, with a spe-
24	cialty or other expertise clinically relevant to the de-
25	vice under review; and

1	"(ii) at least one voting member who is knowl-
2	edgeable about the technology of the device.".
3	(b) Panel Review Process.—Section 513(b)(6) of
4	the Federal Food, Drug, and Cosmetic Act (21 U.S.C.
5	360c(b)(6)) is amended—
6	(1) in subparagraph (A)(iii), by inserting before
7	the period at the end ", including by designating a
8	representative who will be provided a time during the
9	panel meeting to address the panel individually (or
10	accompanied by experts selected by such representa-
11	tive) for the purpose of correcting misstatements of
12	fact or providing clarifying information, subject to
13	the discretion of the panel chairperson"; and
14	(2) by striking subparagraph (B) and inserting
15	the following new subparagraph:
16	" $(B)(i)$ Any meeting of a classification panel for a de-
17	vice that is specifically the subject of review shall—
18	"(I) provide adequate time for initial presen-
19	tations by the person whose device is specifically the
20	subject of a classification panel review and by the
21	Secretary; and
22	"(II) encourage free and open participation by
23	all interested persons.
24	"(ii) Following the initial presentations described in
25	clause (i), the panel may—

1	"(I) pose questions to a designated representative
2	described in subparagraph (A)(iii); and
3	"(II) consider the responses to such questions in
4	the panel's review of the device that is specifically the
5	subject of review by the panel.".
6	SEC. 2227. HUMANITARIAN DEVICE EXEMPTION APPLICA-
7	TION.
8	(a) In General.—Section 520(m) of the Federal
9	Food, Drug, and Cosmetic Act (21 U.S.C. 360j) is amend-
10	ed—
11	(1) in paragraph (1) by striking "fewer than
12	4,000" and inserting "not more than 8,000";
13	(2) in paragraph (2)(A) by striking "fewer than
14	4,000" and inserting "not more than 8,000"; and
15	(3) in paragraph (6)(A)(ii), by striking "4,000"
16	and inserting "8,000"
17	(b) Guidance Document on Probable Benefit.—
18	Not later than 18 months after the date of enactment of
19	this Act, the Secretary of Health and Human Services, act-
20	ing through the Commissioner of Food and Drugs, shall
21	publish a draft guidance document that defines the criteria
22	for establishing "probable benefit" as that term is used in
23	section $520(m)(2)(C)$ of the Federal Food, Drug, and Cos-
24	$metic\ Act\ (21\ U.S.C.\ 360j(m)(2)(C)).$

1	SEC. 2228. CLIA WAIVER STUDY DESIGN GUIDANCE FOR IN
2	VITRO DIAGNOSTICS.
3	(a) Draft Revised Guidance.—Not later than 12
4	months after the date of the enactment of this Act, the Sec-
5	retary of Health and Human Services shall publish a draft
6	guidance that—
7	(1) revises "Section V. Demonstrating Insignifi-
8	cant Risk of an Erroneous Result—'Accuracy'" of the
9	guidance entitled "Recommendations for Clinical
10	Laboratory Improvement Amendments of 1988
11	(CLIA) Waiver Applications for Manufacturers of In
12	Vitro Diagnostic Devices" and dated January 30,
13	2008; and
14	(2) includes guidance on the appropriate use of
15	comparable performance between a waived user and a
16	moderately complex laboratory user to demonstrate
17	accuracy.
18	(b) Final Revised Guidance.—The Secretary of
19	Health and Human Services shall finalize the draft guid-
20	ance published under subsection (a) not later than 12
21	months after the comment period for such draft guidance
22	closes

Subtitle N—Sensible Oversight for Advances Which Technology 2 Regulatory Efficiency 3 4 SEC. 2241. HEALTH SOFTWARE. 5 Section 201 of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 321) is amended by adding at the end the 7 following: 8 "(ss)(1) The term 'health software' means software that does not, through use of an in vitro diagnostic device or signal acquisition system, acquire, process, or analyze an image or physiological signal, is not an accessory, is not 11 an integral part of a device necessary to support the use of the device, is not used in the manufacture and transfusion of blood and blood components to assist in the prevention of disease in humans, and— 16 "(A) is intended for use for administrative or operational support or the processing and mainte-17 18 nance of financial records; "(B) is intended for use in clinical, laboratory, 19 20 or administrative workflow and related recordkeeping; 21 "(C)(i) is intended for use solely in the transfer, 22 aggregation, conversion (in accordance with a present 23 specification), storage, management, retrieval, or 24 transmission of data or information;

1	"(ii) utilizes a connectivity software platform,
2	electronic or electrical hardware, or a physical com-
3	munications infrastructure; and
4	"(iii) is not intended for use—
5	"(I) in active patient monitoring; or
6	"(II) in controlling or altering the functions
7	or parameters of a device that is connected to
8	such software;
9	"(D) is intended for use to organize and present
10	information for health or wellness education or for
11	use in maintaining a healthy lifestyle, including
12	medication adherence and health management tools;
13	"(E) is intended for use to analyze information
14	to provide general health information that does not
15	include patient-specific recommended options to con-
16	sider in the prevention, diagnosis, treatment, cure, or
17	mitigation of a particular disease or condition; or
18	"(F) is intended for use to analyze information
19	to provide patient-specific recommended options to
20	consider in the prevention, diagnosis, treatment, cure,
21	or mitigation of a particular disease or condition.
22	"(2) The term 'accessory' means a product that—
23	"(A) is intended for use with one or more parent
24	devices;

1	"(B) is intended to support, supplement, or aug-
2	ment the performance of one or more parent devices;
3	and
4	"(C) shall be classified by the Secretary—
5	"(i) according to its intended use; and
6	"(ii) independently of any classification of
7	any parent device with which it is used.".
8	SEC. 2242. APPLICABILITY AND INAPPLICABILITY OF REGU-
9	LATION.
10	Subchapter A of chapter V of the Federal Food, Drug,
11	and Cosmetic Act (21 U.S.C. 351 et seq.), as amended by
12	section 2221(a), is further amended by adding at the end
13	the following:
14	"SEC. 524C. HEALTH SOFTWARE.
15	"(a) Inapplicability of Regulation to Health
16	Software.—Except as provided in subsection (b), health
17	software shall not be subject to regulation under this Act.
18	"(b) Exception.—
19	"(1) In general.—Subsection (a) shall not
20	apply with respect to a software product—
21	"(A) of a type described in subparagraph
22	(F) of section $201(ss)(1)$; and
23	"(B) that the Secretary determines poses a
24	significant risk to patient safety.

1	"(2) Considerations.—In making a deter-
2	mination under subparagraph (B) of paragraph (1)
3	with respect to a product to which such paragraph
4	applies, the Secretary shall consider the following:
5	"(A) The likelihood and severity of patient
6	harm if the product were to not perform as in-
7	tended.
8	"(B) The extent to which the product is in-
9	tended to support the clinical judgment of a
10	medical professional.
11	"(C) Whether there is a reasonable oppor-
12	tunity for a medical professional to review the
13	basis of the information or treatment rec-
14	ommendation provided by the product.
15	"(D) The intended user and user environ-
16	ment, such as whether a medical professional
17	will use a software product of a type described
18	in subparagraph (F) of section $201(ss)(1)$.
19	"(c) Delegation.—The Secretary shall delegate pri-
20	mary jurisdiction for regulating a software product deter-
21	mined under subsection (b) to be subject to regulation under
22	this Act to the center at the Food and Drug Administration
23	charged with regulating devices.
24	"(d) Regulation of Software.—

1	"(1) In General.—The Secretary shall review
2	existing regulations and guidance regarding the regu-
3	lation of software under this Act. The Secretary may
4	implement a new framework for the regulation of soft-
5	ware and shall, as appropriate, modify such regula-
6	tions and guidance or issue new regulations or guid-
7	ance.
8	"(2) Issuance by order.—Notwithstanding
9	subchapter II of chapter 5 of title 5, United States
10	Code, the Secretary may modify or issue regulations
11	for the regulation of software under this Act by ad-
12	ministrative order published in the Federal Register
13	following the publication of a proposed order.
14	"(3) Areas under review.—The review of ex-
15	isting regulations and guidance under paragraph (1)
16	may include review of the following areas:
17	"(A) Classification of software.
18	"(B) Standards for development of software.
19	"(C) Standards for validation and
20	verification of software.
21	"(D) Review of software.
22	$``(E)\ Modifications\ to\ software.$
23	"(F) Manufacturing of software.
24	"(G) Quality systems for software.
25	"(H) Labeling requirements for software.

1	"(I) Postmarketing requirements for report-
2	ing of adverse events.
3	"(4) Process for issuing proposed regula-
4	Tions, administrative order, and guidance.—Not
5	later than 18 months after the date of enactment of
6	this section, the Secretary shall consult with external
7	stakeholders (including patients, industry, health care
8	providers, academia, and government) to gather input
9	before issuing regulations, an administrative order,
10	and guidance under this subsection.
11	"(e) Rule of Construction.—Nothing in this sec-
12	tion shall be construed as providing the Secretary with the
13	authority to regulate under this Act any health software
14	product of the type described in subparagraph (F) of section
15	201(ss)(1) unless and until the Secretary has made a deter-
16	$mination\ described\ in\ subsection\ (b)(1)(B)\ with\ respect\ to$
17	such product.".
18	SEC. 2243. EXCLUSION FROM DEFINITION OF DEVICE.
19	Section 201(h) of the Federal Food, Drug, and Cos-
20	metic Act (21 U.S.C. 321) is amended—
21	(1) in subparagraph (2), by striking "or" after
22	"or other animals,";
23	(2) in subparagraph (3), by striking "and" and
24	inserting "or"; and

1	(3) by inserting after subparagraph (3) the fol-
2	lowing:
3	"(4) not health software (other than software de-
4	termined to be a risk to patient safety under section
5	524B(b)), and".
6	Subtitle O—Streamlining Clinical
7	Trials
8	SEC. 2261. PROTECTION OF HUMAN SUBJECTS IN RE-
9	SEARCH; APPLICABILITY OF RULES.
10	(a) In General.—In order to simplify and facilitate
11	compliance by researchers with applicable regulations for
12	the protection of human subjects in research, the Secretary
13	of Health and Human Services shall, to the extent possible
14	and consistent with other statutory provisions, harmonize
15	differences between the HHS Human Subject Regulations
16	and the FDA Human Subject Regulations in accordance
17	with subsection (b).
18	(b) Avoiding Regulatory Duplication and Un-
19	NECESSARY DELAYS.—
20	(1) In General.—The Secretary shall—
21	(A) make such modifications to the provi-
22	sions of the HHS Human Subject Regulations,
23	the FDA Human Subject Regulations, and the
24	vulnerable-populations rules as may be nec-
25	essary—

1	(i) to reduce regulatory duplication
2	and unnecessary delays;
3	(ii) to modernize such provisions in the
4	context of multisite and cooperative research
5	projects; and
6	(iii) to incorporate local consider-
7	ations, community values, and mechanisms
8	to protect vulnerable populations; and
9	(B) ensure that human subject research that
10	is subject to the HHS Human Subject Regula-
11	tions or to the FDA Human Subject Regulations
12	may—
13	(i) use joint or shared review;
14	(ii) rely upon the review of—
15	(I) an independent institutional
16	review board; or
17	(II) an institutional review board
18	of an entity other than the sponsor of
19	the research; or
20	(iii) use similar arrangements to avoid
21	duplication of effort.
22	(2) Regulations and Guidance.—Not later
23	than 36 months after the date of enactment of this
24	Act, the Secretary, acting through the relevant agen-
25	cies and offices of the Department of Health and

1	Human Services, including the Office for Human Re-
2	search Protections and relevant agencies and offices of
3	the Food and Drug Administration, shall issue such
4	regulations and guidance and take such other actions
5	as may be necessary to implement this section and
6	help to facilitate the broader use of single, central, or
7	lead institutional review boards. Such regulations
8	and guidance shall clarify the requirements and poli-
9	cies relating to the following:
10	(A) Arrangements to avoid duplication de-
11	$scribed\ in\ paragraph\ (1)(A)(i),\ including$ —
12	(i) delineating the roles of institutional
13	review boards in multisite or cooperative,
14	multisite studies where one or more local
15	institutional review boards are relied upon,
16	or similar arrangements are used;
17	(ii) the risks and benefits to human
18	subjects;
19	(iii) standardizing the informed con-
20	sent and other processes and legal docu-
21	ments; and
22	(iv) incorporating community values
23	through the use of local institutional review
24	boards while continuing to use central or
25	lead institutional review boards.

1	(B) Concerns about regulatory and legal li-
2	ability contributing to decisions by the sponsors
3	of research to rely on local institutional review
4	boards for multisite research.
5	(3) Consultation.—In issuing regulations or
6	guidance under paragraph (2), the Secretary shall
7	consult with stakeholders (including researchers, aca-
8	demic organizations, hospitals, institutional research
9	boards, pharmaceutical, biotechnology and medical
10	device developers, clinical research organizations, pa-
11	tient groups, and others).
12	(c) Timing.—The Secretary shall complete the harmo-
13	nization described in subsection (a) not later than 36
14	months after the date of enactment of this Act.
15	(d) Progress Report.—Not later than 24 months
16	after the date of enactment of this Act, the Secretary shall
17	submit to Congress a report on the progress made toward
18	completing such harmonization.
19	(e) Draft NIH Policy.—Not later than 12 months
20	after the date of enactment of this Act, the Secretary, acting
21	through the Director of the National Institutes of Health,
22	shall finalize the draft policy entitled "Draft NIH Policy
23	on Use of a Single Institutional Review Board for Multi-
24	Site Research".

1	(1) Human subject regulations.—In this
2	section:
3	(A) FDA HUMAN SUBJECT REGULATIONS.—
4	The term "FDA Human Subject Regulations"
5	means the provisions of parts 50, 56, 312, and
6	812 of title 21, Code of Federal Regulations (or
7	any successor regulations).
8	(B) HHS HUMAN SUBJECT REGULA-
9	Tions.—The term "HHS Human Subject Regu-
10	lations" means the provisions of subpart A of
11	part 46 of title 45, Code of Federal Regulations
12	(or any successor regulations).
13	(C) Vulnerable-populations rules.—
14	The term "vulnerable-populations rules"—
15	(i) subject to clause (ii), means the
16	provisions of subparts B through D of such
17	part 46 (or any successor regulations); or
18	(ii) as applicable to research that is
19	subject to the FDA Human Subject Regula-
20	tions, means the provisions applicable to
21	vulnerable populations under part 56 of
22	such title 21 (or any successor regulations)
23	and subpart D of part 50 of such title 21
24	(or any successor regulations).
25	(2) Other definitions.—In this section:

1	(A) Institutional review board.—The
2	term "institutional review board" has the mean-
3	ing that applies to the term "institutional review
4	board" under the HHS Human Subject Regula-
5	tions.
6	(B) Lead institutional review
7	BOARD.—The term "lead institutional review
8	board" means an institutional review board than
9	otherwise meets the requirements of the HHS
10	Human Subject Regulations and enters into a
11	written agreement with an institution, another
12	institutional review board, a sponsor, or a prin-
13	cipal investigator to approve and oversee human
14	subject research that is conducted at multiple lo-
15	cations. References to an institutional review
16	board include an institutional review board than
17	serves a single institution as well as a lead insti-
18	tutional review board.
19	SEC. 2262. USE OF NON-LOCAL INSTITUTIONAL REVIEW
20	BOARDS FOR REVIEW OF INVESTIGATIONAL
21	DEVICE EXEMPTIONS AND HUMAN DEVICE
22	EXEMPTIONS.
23	(a) In General.—Section 520 of the Federal Food,
24	Drug, and Cosmetic Act (21 U.S.C. 360(j)) is amended—
25	(1) in subsection $(g)(3)$ —

1	(A) by striking "local" each place it ap-
2	pears; and
3	(B) in subparagraph $(A)(i)$, by striking
4	"which has been"; and
5	(2) in subsection $(m)(4)$ —
6	(A) by striking "local" each place it ap-
7	pears; and
8	(B) by striking subparagraph (A) and in-
9	serting the following new subparagraph:
10	"(A) in facilities in which clinical testing of de-
11	vices is supervised by an institutional review com-
12	mittee established in accordance with the regulations
13	of the Secretary, and".
14	(b) Regulations.—Not later than 12 months after the
15	date of the enactment of this Act, the Secretary of Health
16	and Human Services shall revise or issue such regulations
17	or guidance as may be necessary to carry out the amend-
18	ments made by subsection (a).
19	SEC. 2263. ALTERATION OR WAIVER OF INFORMED CON-
20	SENT FOR CLINICAL INVESTIGATIONS.
21	(a) Devices.—Section 520(g)(3) of the Federal Food,
22	Drug, and Cosmetic Act (21 U.S.C. 360j(g)(3)) is amend-
23	ed—
24	(1) in subparagraph (D), by striking "except
25	where subject to such conditions as the Secretary may

1	prescribe, the investigator" and inserting the fol-
2	lowing: "except where, subject to such conditions as
3	the Secretary may prescribe—
4	"(i) the proposed clinical testing poses no
5	more than minimal risk to the human subject
6	and includes appropriate safeguards to protect
7	the rights, safety, and welfare of the human sub-
8	$ject;\ or$
9	"(ii) the investigator"; and
10	(2) in the matter following subparagraph (D), by
11	striking "subparagraph (D)" and inserting "subpara-
12	$graph\ (D)(ii)$ ".
13	(b) DRUGS.—Section 505(i)(4) of the Federal Food,
14	Drug, and Cosmetic Act (21 U.S.C. 355(i)(4)) is amended
15	by striking "except where it is not feasible or it is contrary
16	to the best interests of such human beings" and inserting
17	"except where it is not feasible, it is contrary to the best
18	interests of such human beings, or the proposed clinical test-
19	ing poses no more than minimal risk to such human beings
20	and includes appropriate safeguards as prescribed to pro-
21	tect the rights, safety, and welfare of such human beings".

1	Subtitle P—Improving Scientific
2	Expertise and Outreach at FDA
3	SEC. 2281. SILVIO O. CONTE SENIOR BIOMEDICAL RE-
4	SEARCH SERVICE.
5	(a) Hiring and Retention Authority.—Section
6	228 of the Public Health Service Act (42 U.S.C. 237) is
7	amended—
8	(1) in the section heading, by inserting "AND
9	BIOMEDICAL PRODUCT ASSESSMENT" after "RE-
10	SEARCH'';
11	(2) in subsection (a)(1), by striking "Silvio O.
12	Conte Senior Biomedical Research Service, not to ex-
13	ceed 500 members" and inserting "Silvio O. Conte
14	Senior Biomedical Research and Biomedical Product
15	Assessment Service (in this section referred to as the
16	'Service'), the purpose of which is to recruit and re-
17	tain competitive and qualified scientific and technical
18	experts outstanding in the field of biomedical re-
19	search, clinical research evaluation, and biomedical
20	product assessment";
21	(3) by amending subsection (a)(2) to read as fol-
22	lows:
23	"(2) The authority established in paragraph (1) may
24	not be construed to require the Secretary to reduce the num-
25	ber of employees serving under any other employment sys-

1	tem in order to offset the number of members serving in
2	the Service.";
3	(4) in subsection (b)—
4	(A) in the matter preceding paragraph (1),
5	by striking "or clinical research evaluation" and
6	inserting ", clinical research evaluation or bio-
7	medical product assessment"; and
8	(B) in paragraph (1), by inserting "or a
9	master's level degree in engineering,
10	bioinformatics, or a related or emerging field,"
11	after the comma;
12	(5) in subsection $(d)(2)$, by striking "and shall
13	not exceed the rate payable for level I of the Executive
14	Schedule unless approved by the President under sec-
15	tion 5377(d)(2) of title 5, United States Code" and
16	inserting "and shall not exceed the rate payable for
17	the President";
18	(6) by striking subsection (e); and
19	(7) by redesignating subsections (f) and (g) as
20	subsections (e) and (f), respectively.
21	(b) Report.—Not later than 3 years after the date
22	of the enactment of this Act, the Secretary of Health and
23	Human Services shall submit, and publish on the website
24	of the Department of Health and Human Services a report
25	on the implementation of the amendments made by sub-

1	section (a), including whether the amendments have im-
2	proved the ability of the Food and Drug Administration
3	to hire and retain qualified experts to fulfill obligations
4	specified under user fee agreements.
5	SEC. 2282. ENABLING FDA SCIENTIFIC ENGAGEMENT.
6	It is the sense of Congress that the participation in,
7	or sponsorship of, scientific conferences and meetings is es-
8	sential to the mission of the Food and Drug Administra-
9	tion.
10	SEC. 2283. REAGAN-UDALL FOUNDATION FOR THE FOOD
11	AND DRUG ADMINISTRATION.
12	(a) Board of Directors.—
13	(1) Composition and size.—Section
14	770(d)(1)(C) of the Federal Food, Drug, and Cosmetic
15	Act (21 U.S.C. 379dd(d)(1)(C)) is amended—
16	(A) by redesignating clause (ii) as clause
17	(iii);
18	(B) by inserting after clause (i) the fol-
19	lowing:
20	"(ii) Additional members.—The
21	Board, through amendments to the bylaws
22	of the Foundation, may provide that the
23	number of voting members of the Board
24	shall be a number (to be specified in such
25	amendment) greater than 14. Any Board

1	positions that are established by any such
2	amendment shall be appointed (by majority
3	vote) by the individuals who, as of the date
4	of such amendment, are voting members of
5	the Board and persons so appointed may
6	represent any of the categories specified in
7	subclauses (I) through (V) of clause (i), so
8	long as no more than 30 percent of the total
9	voting members of the Board (including
10	members whose positions are established by
11	such amendment) are representatives of the
12	general pharmaceutical, device, food, cos-
13	metic, and biotechnology industries."; and
14	(C) in clause $(iii)(I)$, as redesignated by
15	subparagraph (A), by striking "The ex officio
16	members shall ensure" and inserting "The ex
17	officio members, acting pursuant to clause (i),
18	and the Board, acting pursuant to clause (ii),
19	shall ensure".
20	(2) Federal employees allowed to serve
21	ON BOARD.—Clause (iii)(II) of section 770(d)(1)(C) of
22	the Federal Food, Drug, and Cosmetic Act (21 U.S.C.
23	379dd(d)(1)(C)), as redesignated by paragraph

(1)(A), is amended by adding at the end the fol-

lowing: "For purposes of this section, the term 'em-

24

1	ployee of the Federal Government' does not include a
2	'special Government employee', as that term is de-
3	fined in section 202(a) of title 18, United States
4	Code.".
5	(3) Staggered terms.—Subparagraph (A) of
6	section $770(d)(3)$ of the Federal Food, Drug, and Cos-
7	metic Act (21 U.S.C. 379dd(d)(3)) is amended to read
8	as follows:
9	"(A) Term.—The term of office of each
10	member of the Board appointed under para-
11	$graph\ (1)(C)(i),\ and\ the\ term\ of\ office\ of\ any$
12	member of the Board whose position is estab-
13	lished pursuant to paragraph (1)(C)(ii), shall be
14	4 years, except that—
15	"(i) the terms of offices for the members
16	of the Board initially appointed under
17	paragraph (1)(C)(i) shall expire on a stag-
18	gered basis as determined by the ex officio
19	members; and
20	"(ii) the terms of office for the persons
21	initially appointed to positions established
22	pursuant to $paragraph$ (1)(C)(ii) may be
23	made to expire on a staggered basis, as de-
24	termined by the individuals who as of the

1	date of the amendment establishing such po-
2	sitions, are members of the Board.".
3	(b) Executive Director Compensation.—Section
4	770(g)(2) of the Federal Food, Drug, and Cosmetic Act (21
5	$U.S.C.\ 379dd(g)(2))$ is amended by striking 'but shall not
6	be greater than the compensation of the Commissioner".
7	(c) Separation of Funds.—Section 770(m) of the
8	Federal Food, Drug, and Cosmetic Act (21 U.S.C.
9	379dd(m)) is amended by striking "are held in separate
10	accounts from funds received from entities under subsection
11	(i)" and inserting "are managed as individual pro-
12	grammatic funds under subsection (i), according to best ac-
13	counting practices".
14	SEC. 2284. COLLECTION OF CERTAIN VOLUNTARY INFORMA-
15	TION EXEMPTED FROM PAPERWORK REDUC-
16	TION ACT.
17	Chapter VII of the Federal Food, Drug, and Cosmetic
18	Act is amended by inserting after section 708 of such Act
19	(21 U.S.C. 379) the following:
20	"SEC. 708A. COLLECTION OF CERTAIN VOLUNTARY INFOR-
21	MATION EXEMPTED FROM PAPERWORK RE-
22	DUCTION ACT.
23	"Chapter 35 of title 44, United States Code, shall not
24	apply to the collection from patients, industry, academia,
25	and other stakeholders, of voluntary information such as

1 through voluntary surveys or questionnaires, initiated by

2	the Secretary.".			
3	SEC. 2285. HIRING AUTHORITY FOR SCIENTIFIC, TECH-			
4	NICAL, AND PROFESSIONAL PERSONNEL.			
5	(a) In General.—The Federal Food, Drug, and Cos-			
6	metic Act is amended by inserting after section 714 (21			
7	U.S.C. 379d-3) the following:			
8	"SEC. 714A. ADDITIONAL HIRING AUTHORITY.			
9	"(a) In General.—The Secretary may, without re-			
10	gard to the provisions of title 5, United States Code, gov-			
11	erning appointments in the competitive service, appoint			
12	qualified candidates to scientific, technical, or professional			
13	positions within the following centers of the Food and Drug			
14	Administration:			
15	"(1) The Center for Drug Evaluation and Re-			
16	search.			
17	"(2) The Center for Biologics Evaluation and			
18	Research.			
19	"(3) The Center for Devices and Radiological			
20	Health.			
21	Such positions shall be within the competitive service.			
22	"(b) Compensation.—			
23	"(1) In GENERAL.—Notwithstanding any other			
24	provision of law, including any requirement with re-			
25	spect to General Schedule pay rates under subchapter			

1	III of chapter 53 of title 5, United States Code, and
2	consistent with the requirements of paragraph (2), the
3	Secretary may determine and fix—
4	"(A) the annual rate of pay of any indi-
5	vidual appointed under subsection (a); and
6	"(B) for purposes of retaining qualified em-
7	ployees, the annual rate of pay for any highly
8	qualified scientific, technical, or professional per-
9	sonnel appointed to a position at any of the cen-
10	ters listed under subsection (a) before the date of
11	enactment of this section.
12	"(2) Limitation.—The annual rate of pay es-
13	tablished pursuant to paragraph (1) may not exceed
14	the annual rate of pay of the President.
15	"(c) Sunset.—The authority to appoint employees
16	under this section shall terminate on September 30, 2022.
17	"(d) Report.—
18	"(1) In general.—Not later than September 30,
19	2021, the Secretary shall submit a report to Congress
20	that examines the extent to which the authority to ap-
21	point and retain personnel under this section en-
22	hanced the Food and Drug Administration's ability
23	to meet the agency's critical need for highly qualified
24	individuals for scientific, technical, or professional
25	positions.

1	"(2) RECOMMENDATIONS.—The report under				
2	paragraph (1) shall include the recommendations of				
3	the Secretary on—				
4	"(A) whether the authority to appoint per-				
5	sonnel under this section should be reauthorized;				
6	and				
7	"(B) other personnel authorities that would				
8	help the Food and Drug Administration to better				
9	recruit and retain highly qualified individuals				
10	for scientific, technical, or professional positions				
11	in the agency's medical product centers.".				
12	(b) Rule of Construction.—The authority provided				
13	by section 714A of the Federal Food, Drug, and Cosmetic				
14	Act (as added by subsection (a)) shall not be construed to				
15	affect the authority provided under section 714 of such Act.				
16	Subtitle Q—Exempting From				
17	Sequestration Certain User Fees				
18	SEC. 2301. EXEMPTING FROM SEQUESTRATION CERTAIN				
19	USER FEES OF FOOD AND DRUG ADMINISTRA-				
20	TION.				
21	The Balanced Budget and Emergency Deficit Control				
22	Act of 1985 is amended—				
23	(1) in section $255(g)(1)(A)$ (2 U.S.C.				
24	905(g)(1)(A)), by inserting after the item relating to				
25	"Financial Agent Services" the following new item:				

1	"Food and Drug Administration, Salaries			
2	and Expenses, but only the portion of appropria-			
3	tions under such account corresponding to fees			
4	collected under sections 736, 738, 740, 741			
5	744B, and 744H of the Federal Food, Drug, and			
6	Cosmetic Act (75–9911–0–1–554)."; and			
7	(2) in section 256(h) (2 U.S.C. 906(h)), by add-			
8	ing at the end the following new paragraph:			
9	"(5) Notwithstanding any other provision of law,			
10	this subsection shall not apply with respect to the por-			
11	tion of administrative expenses incurred by the Food			
12	and Drug Administration that are funded through			
13	fees collected under sections 736, 738, 740, 741, 744B,			
14	and 744H of the Federal Food, Drug, and Cosmetic			
15	Act.".			
16	TITLE III—DELIVERY			
17	Subtitle A—Interoperability			
18	SEC. 3001. ENSURING INTEROPERABILITY OF HEALTH IN-			
19	FORMATION TECHNOLOGY.			
20	(a) Interoperability Standards.—			
21	(1) In general.—Subtitle A of title XXX of the			
22	Public Health Service Act (42 U.S.C. 300jj–11 et seq.)			
23	is amended by adding at the end the following new			
24	section:			

1	"SEC. 3010. ENSURING INTEROPERABILITY OF HEALTH IN-
2	FORMATION TECHNOLOGY.
3	"(a) Interoperability.—In order for health infor-
4	mation technology to be considered interoperable, such tech-
5	nology must satisfy the following criteria:
6	"(1) Secure transfer.—The technology allows
7	the secure transfer of the entirety of a patient's data
8	from any and all health information technology for
9	authorized use under applicable law.
10	"(2) Complete access to health data.—The
11	technology allows access to the entirety of a patient's
12	available data for authorized use under applicable
13	law without special effort, as defined by recommenda-
14	tions for interoperability standards adopted under
15	section 3004, by the requestor of such data unless such
16	data is not disclosable under applicable law.
17	"(3) No information blocking.—The tech-
18	nology is not configured, set up, or implemented to
19	engage in information blocking, as defined in section
20	3010A(f).
21	"(b) Categories for Interoperability Stand-
22	ARDS.—The categories described in this subsection, with re-
23	spect to standards for determining if health information
24	technology is interoperable, consistent with the criteria de-
25	scribed in subsection (a), include the following categories
26	of standards:

1	"(1) Standards with respect to vocabulary and
2	terminology.
3	"(2) Standards with respect to content and
4	structure.
5	"(3) Standards with respect to transport of in-
6	formation.
7	"(4) Security standards.
8	"(5) Service standards.".
9	(2) Guidance.—Not later than January 1,
10	2017, the Secretary of Health and Human Services,
11	through the National Coordinator of the Office of the
12	National Coordinator for Health Information Tech-
13	nology, shall issue guidance with respect to the imple-
14	mentation of section 3010 of the Public Health Serv-
15	ice Act, as added by paragraph (1), including with
16	respect to defining and providing examples of author-
17	ized use of health information technology, as described
18	in such section.
19	(b) Improvements to Recommendation Proc-
20	ESS.—
21	(1) HIT POLICY COMMITTEE TO INCORPORATE
22	POLICIES FOR UPDATES TO INTEROPERABILITY
23	STANDARDS.—Section 3002 of the Public Health
24	Service Act (42 U.S.C. 300jj-12) is amended—
25	(A) in subsection (a) —

1	(i) by striking "National Coordinator"
2	and inserting "Secretary, in consultation
3	with the National Coordinator,"; and
4	(ii) by adding at the end the following
5	new sentence: "The HIT Policy Committee
6	is authorized only to provide policy and
7	priority recommendations to the Secretary
8	and not authorized to otherwise affect the
9	development or modification of any stand-
10	ard, implementation specification, or cer-
11	tification criterion under this title."; and
12	(B) in subsection $(b)(2)$ —
13	(i) in subparagraph (A), in the first
14	sentence—
15	(I) by striking "The HIT Policy
16	Committee" and inserting "Subject to
17	subparagraph (D), the HIT Policy
18	Committee"; and
19	(II) by inserting "(including the
20	areas in which modifications and ad-
21	ditions to interoperability standards
22	under section 3010 are needed for the
23	electronic exchange and use of health
24	information for purposes of adoption of
25	such modifications and additions

1	under section 3004)" after "section
2	3004".
3	(ii) by adding at the end the following
4	new subparagraph:
5	"(D) Special rule related to inter-
6	OPERABILITY.—Any recommendation made by
7	the HIT Policy Committee on or after the date
8	of the enactment of this subparagraph with re-
9	spect to interoperability of health information
10	technology shall be consistent with the criteria
11	described in subsection (a) of section 3010.".
12	(2) Sunset of hit standards committee.—
13	Section 3003 of the Public Health Service Act (42
14	U.S.C. 300jj-13) is amended by adding at the end the
15	following new subsection:
16	"(f) Termination.—The HIT Standards Committee
17	shall terminate on the date that is 90 days after the date
18	of the enactment of this subsection.".
19	(3) Standards development organiza-
20	Tions.—Title XXX of the Public Health Service Act
21	is amended by inserting after section 3003 the fol-
22	lowing new section:

1	"SEC. 3003A.	RECOMMENDATIONS	FOR	STANDARDS
2	THE	ROUGH CONTRACTS W	ITH ST	ANDARDS DE-
3	VEL	OPMENT ORGANIZATI	ONS.	
4	"(a) Contra	ACTS.—		
5	"(1) I	N GENERAL.—For pu	ırposes	of activities
6	conducted u	under this title, the S	<i>lecretar</i>	ry shall enter
7	into contrac	ts with health care sto	ındard:	s development
8	organization	ns accredited by the	Ameri	can National
9	Standards 1	Institute to carry out	the du	ties described
10	in subsection	n (b), as applicable.		
11	"(2) Ti	IMING FOR FIRST CON	TRACT.	—As soon as
12	practicable	after the date of the en	nactme	nt of this sec-
13	tion, the Se	cretary shall enter in	to the	first contract
14	under parag	graph (1).		
15	"(3) I	PERIOD OF CONTRAC	∑ T.— E	ach contract
16	under parag	graph (1) shall be for	a perio	od determined
17	necessary by	y the Secretary, in co	msulta	tion with the
18	National Co	ordinator, to carry ou	it the a	applicable du-
19	ties describe	d in subsection (b).		
20	"(4) A	PPROPRIATE ORGANIZ	ZATION	s.—The Sec-
21	retary shall	t ensure the most ap	propri	ate organiza-
22	tions describ	bed in paragraph (1)	are sel	ected for each
23	contract und	der such paragraph.		
24	"(5) A	LLOWANCE FOR VARL	ATIONS	.—Standards
25	developed p	oursuant to a contra	ect une	der this sub-
26	section, and	the methods to test s	uch ste	andards, shall

1 allow for variations on such standards as long as 2 such variations are consistent with the standards so developed under this section. 3 "(b) Duties.— 4 "(1) Initial contract.—Under the initial con-5 6 tract under subsection (a)(1), the standards develop-7 ment organizations— 8 "(A) shall provide to the Secretary, in con-9 sultation with the National Coordinator, for adoption under section 3004, recommendations, 10 in accordance with section 3010, for interoper-11 12 ability standards, and methods to test such 13 standards, consistent with the criteria described 14 in subsection (a) of such section and with respect 15 to the categories described in subsection (b)(1) of such section; and 16 17 "(B) may provide to the Secretary rec-18 ommendations described in paragraph (2).

"(2) Subsequent contract, the organizations shall provide to the Secretary, in consultation with the National Coordinator, for adoption under section 3004 recommendations for any standards (including interoperability standards and methods to test such standards), implementation specifications, and certifi-

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- cation criteria (and modifications, including additions, to such standards, specifications, and criteria), which are in accordance with the policies and priorities developed by the Secretary, in consultation with the National Coordinator.
 - "(3) MULTIPLE METHODS TO TEST INTEROPER-ABILITY STANDARDS.—For the purposes of developing methods to test interoperability standards for adoption under section 3004, the Secretary shall ensure that contracts under this section allow for multiple methods to test such standards to account for variations in the adoption of such standards that do not conflict with section 3010(a).
 - "(c) Modifications and Subsequent Contracts.—
 - "(1) IN GENERAL.—The Secretary, in consultation with the National Coordinator, shall periodically conduct hearings to evaluate and review the standards, implementation specifications, and certification criteria adopted under section 3004 for purposes of determining if modifications, including any additions, are needed with respect to such standards, specifications, and criteria.
 - "(2) Contract trigger.—Based on the needs for standards, implementation specifications, and certification criteria (and modifications, including addi-

1	tions, to such standards, specifications, and criteria)
2	under this title, as determined by the Secretary, in
3	consultation with the National Coordinator, the Sec-
4	retary shall, as needed, enter into contracts under
5	subsection (a) in addition to the initial contract.
6	"(d) Authorization of Appropriations.—There is
7	authorized to be appropriated \$10,000,000 for contracts
8	under subsection (a), to remain available until expended.".
9	(4) Modifications to role of onchit.—Sec-
10	tion 3001(c)(1)(A) of the Public Health Service Act
11	(42 U.S.C. 300jj-11(c)(1)(A)) is amended by insert-
12	ing "for recommendations made before the date of the
13	enactment of the 21st Century Cures Act," before "re-
14	view and determine".
15	(c) Adoption.—Section 3004 of the Public Health
16	Service Act (42 U.S.C. 300jj–14) is amended—
17	(1) in subsection (a)—
18	(A) in paragraph (1), by inserting after
19	"section 3001(c)" the following: "(or, subject to
20	subsection (c), in the case of a standard, speci-
21	fication, or criterion recommended on or after
22	the date of the enactment of the 21st Century
23	Cures Act, after the date of submission of the rec-
24	ommendation to the Secretary under section
25	3003A)"; and

1	(B) in paragraph (2)(B), by striking "and
2	the HIT Standards Committee";
3	(2) in subsection (b), by adding at the end the
4	following new paragraph:
5	"(4) Limitation.—The Secretary may not adopt
6	any standards, implementation specifications, or cer-
7	tification criteria under this subsection or subsection
8	(a) that are inconsistent with or duplicative of an
9	interoperability standard adopted under this section,
10	in accordance with subsections (c) and (d). In the
11	case of a standard, specification, or criterion that has
12	been adopted under this section and is inconsistent or
13	duplicative of such an interoperability standard than
14	is subsequently adopted under this section, such inter-
15	operability standard shall supercede such other stand-
16	ard, specification, or criterion and such other stand-
17	ard, specification, or criterion shall no longer be con-
18	sidered adopted under this section beginning on the
19	date that such interoperability standard becomes effec-
20	tive."; and
21	(3) by adding at the end the following new sub-
22	sections:
23	"(c) Adoption of Initial Interoperability Stand-
24	ARDS.—Notwithstanding the previous subsections of this
25	section, the following shall apply in the case of the initial

1	set of interoperability standards recommended under sec-
2	tion 3003A:
3	"(1) Review of Standards.—Not later than 90
4	days after the date of receipt of recommendations for
5	such interoperability standards, the Secretary, in con-
6	sultation with the National Coordinator and rep-
7	resentatives of other relevant Federal agencies, shall
8	jointly review such standards and shall determine
9	whether or not to propose adoption of such standards.
10	"(2) Determination to Adopt.—If the Sec-
11	retary determines—
12	"(A) to propose adoption of such standards,
13	the Secretary shall, by regulation under section
14	553 of title 5, United States Code, determine
15	whether or not to adopt such standards; or
16	"(B) not to propose adoption of such stand-
17	ards, the Secretary shall notify the applicable
18	standards development organizations with a con-
19	tract under section 3003A in writing of such de-
20	termination and the reasons for not proposing
21	the adoption of the recommendation for such
22	standards.
23	"(3) Publication.—The Secretary shall provide
24	for publication in the Federal Register of all deter-

1	minations made by the Secretary under paragraph
2	(1).
3	"(4) APPLICATION.—Any standard adopted
4	under this subsection shall be effective 12 months after
5	the date of publication of the determination to adopt
6	such standard.
7	"(d) Rules for Adoption.—In the case of a stand-
8	ard (including interoperability standard), implementation
9	specification, or certification criteria adopted under this
10	section on or after the date of the enactment of the 21st
11	Century Cures Act, the following shall apply:
12	"(1) In general.—Except as provided in para-
13	graph (2), any such standard (including interoper-
14	ability standard), implementation specification, or
15	certification criterion shall be a standard, specifica-
16	tion, or criterion that has been recommended by the
17	standards development organizations with which the
18	Secretary has entered into a contract under section
19	3003A.
20	"(2) Special rule if no standard, specifica-
21	TION, OR CRITERION RECOMMENDED.—If no standard
22	is recommended under paragraph (1)—
23	"(A) in the case of interoperability stand-
24	ards, relating to a category described in section
25	<i>3010(b)</i> —

1	"(i) paragraph (1) shall not apply;
2	and
3	"(ii) paragraph (4) shall apply; or
4	"(B) in the case of any other standard, im-
5	plementation specification, or certification cri-
6	teria, relating to a policy or priority to carry
7	out this title, as determined by the Secretary, in
8	consultation with the National Coordinator—
9	"(i) paragraph (1) shall not apply;
10	and
11	"(ii) paragraph (4) shall apply.
12	"(3) Effective date.—Any standard, imple-
13	mentation specification, or certification criterion
14	adopted under this section shall be effective 12 months
15	after the date of publication of the final rule to adopt
16	such standard, implementation specification, or cer-
17	tification criterion.
18	"(4) Assistance to the secretary.—In com-
19	plying with the requirements of this subsection, the
20	Secretary shall rely on the recommendations of the
21	National Committee on Vital and Health Statistics
22	established under section 306(k), and shall consult
23	with appropriate Federal and State agencies and pri-
24	vate organizations. The Secretary shall publish in the
25	Federal Register any recommendation of the National

1	Committee on Vital and Health Statistics regarding
2	the adoption of a standard, implementation specifica-
3	tion, or certification criterion under this section. Any
4	standard, implementation specification, or certifi-
5	cation criterion adopted pursuant to this paragraph
6	shall be promulgated in accordance with the rule-
7	making procedures of subchapter III of chapter 5 of
8	title 5, United States Code.".
9	(d) Reports and Notifications.—Section 3010 of
10	the Public Health Service Act, as added by subsection (a),
11	is amended by adding at the end the following new sub-
12	section:
13	"(c) Dissemination of Information.—
14	"(1) Initial summary report.—Not later than
15	July 1, 2017, the Secretary, after consultation with
16	relevant stakeholders, shall submit to Congress and
17	provide for publication in the Federal Register and
18	the posting on the Internet website of the Office of the
19	National Coordinator for Health Information Tech-
20	nology of a report on the following:
21	"(A) The initial set of interoperability
22	$standards\ adopted\ under\ section\ 3004 (c).$
23	"(B) The strategies for achieving wide-
24	$spread\ interoperability.$

1	"(C) An overview of the extent to which
2	electronic health records and health information
3	technology offered as of such date satisfy such
4	$initial\ set.$
5	"(D) Any barriers that are preventing wide-
6	$spread\ interoperability.$
7	"(E) The plan and milestones, including
8	specific steps, to achieve widespread interoper-
9	ability.
10	"(2) Followup Determination and Report
11	on widespread interoperability.—Not later than
12	December 31, 2019, the Secretary shall provide for
13	publication in the Federal Register and the posting
14	on the Internet website of the Office of the National
15	Coordinator for Health Information Technology of the
16	following:
17	"(A) A determination by the Secretary
18	whether the goal of widespread interoperability
19	has been achieved.
20	"(B) A list identifying the vendors of, or
21	other entities offering, qualified electronic health
22	records, which categorizes such entities, with re-
23	spect to such records, as in compliance or not in
24	compliance with the certification criteria de-
25	scribed in section 3001(c)(5)(B)(ii) and with the

1	requirements	under	clause	(i)	of	section
2	3001(c)(5)(C)	(includi	ng with	the	terms	of the
3	attestation an	nd other	requirer	nents	s unde	er such
1	clause).					

- "(C) Actions that may be taken by entities identified under subparagraph (B) as not being in compliance with such criteria and requirements in order for such entities to become in compliance with such criteria and requirements.
- "(D) **Penalties** described insection 3010A(d) to which entities, with respect to such qualified electronic health records, beginning January 1, 2019, are subject if such technology and entities are not in compliance with the certification criteriadescribedinsection 3001(c)(5)(B)(ii) and with the requirements under clause (i) of section 3001(c)(5)(C), respectively.
- "(3) Ongoing publication of recommendations.—The Secretary shall provide for publication in the Federal Register and the posting on the Internet website of the Office of the National Coordinator for Health Information Technology of all recommendations made under this section."

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1	(e) Certification and Other Enforcement Provi-
2	SIONS.—
3	(1) Certification of qualified electronic
4	HEALTH RECORDS.—
5	(A) In General.—Section 3007(b) of the
6	Public Health Service Act (42 U.S.C. 300jj-
7	17(b)) is amended by striking "under section
8	3001(c)(3) to be in compliance with" and all
9	that follows through the period at the end and
10	inserting "under section $3001(c)(3)$ —
11	"(1) for certifications made before January 1,
12	2018, to be in compliance with applicable standards
13	adopted under subsections (a) and (b) of section 3004;
14	and
15	"(2) for certifications made on or after January
16	1, 2018, to be in compliance with applicable stand-
17	ards adopted under subsections (a) and (b) of section
18	3004 and to be interoperable in accordance with sec-
19	tion 3010, including by being in compliance with
20	interoperability standards adopted under section
21	3004.".
22	(B) Requirements of Secretary.—Sec-
23	tion 3001(c)(5) of the Public Health Service Act
24	(42 U.S.C. 300jj-11(c)(5)) is amended—

1	(i) by amending subparagraph (B) of
2	such section to read as follows:
3	"(B) Certification criteria de-
4	SCRIBED.—In this title, the term 'certification
5	criteria' means, with respect to qualified elec-
6	tronic health records—
7	"(i) for certifications made before Jan-
8	uary 1, 2018, criteria to establish that the
9	records meet standards and implementation
10	specifications adopted under subsections (a)
11	and (b) of section 3004 for qualified elec-
12	tronic health records; and
13	"(ii) for certifications made on or after
14	January 1, 2018, criteria described in
15	clause (i) and criteria to establish that the
16	records are interoperable, in accordance
17	with section 3010, including by being in
18	compliance with interoperability standards
19	adopted under section 3004."; and
20	(ii) by adding at the end the following
21	new subparagraph:
22	"(C) Enforcement; decertifications.—
23	"(i) Requirements.—Under any pro-
24	gram kept or recognized under subpara-
25	graph (A), the Secretary shall ensure that

1	any vendor of or other entity offering quali-
2	fied electronic health records seeking a cer-
3	tification of such records under such pro-
4	gram on or after January 1, 2018, shall, as
5	a condition of certification (and mainte-
6	nance of certification) of such a record
7	under such program—
8	"(I) provide to the Secretary an
9	attestation—
10	"(aa) that the entity, unless
11	for a legitimate purpose specified
12	by the Secretary, has not taken
13	any action, including through
14	any financial, administrative, or
15	technological barrier, which the
16	entity knows or should know (as
17	defined in section $1128A(i)(7)$ of
18	the Social Security Act), is to
19	limit or restrict the exchange of
20	information or to prevent or
21	disincentivize widespread inter-
22	operability between any providers
23	using such records or other health
24	information technology in connec-
25	tion with such record:

1	"(bb) on the pricing informa-
2	tion described in clause (v) for
3	purposes of the portal created
4	under paragraph (9); that such
5	information will be available on a
6	public Web site of such entity and
7	in marketing materials, commu-
8	nications statements, and other
9	assertions of such entity related to
10	such record; and that the entity
11	will voluntarily provide such in-
12	formation to customers prior to
13	providing any qualified electronic
14	health records or related product
15	or service (including subsequent
16	updates, add-ons, or additional
17	products or services to be provided
18	during the course of an on-going
19	contract), prospective customers
20	(such as persons who request or
21	receive a quotation, estimate, or
22	other similar marketing or pro-
23	motional material), and other
24	persons who request such informa-
25	tion;

1	"(cc) that the software with
2	respect to such records have pub-
3	lished application programming
4	interfaces for medical records
5	data, search and indexing, seman-
6	tic harmonization and vocabulary
7	translation, and user interface ap-
8	plications;
9	"(dd) that the entity has suc-
10	cessfully tested the use of the
11	record in the type of setting in
12	which it would be marketed;
13	"(ee) the entity has in place
14	implementation guidelines for
15	such record that support inter-
16	operability, consistent with sec-
17	tion 3010; and
18	"(ff) that the entity has in
19	place data sharing programs or
20	capabilities based on common
21	data elements through application
22	programming interfaces without
23	the requirement for vendor-specific
24	interfaces;

"(II) publish applicat	tion pro-
gramming interfaces and	associated
documentation, with respect	to such
records, for medical recor	rds data,
search and indexing, semant	ic harmo-
nization and vocabulary tr	ranslation,
and user interface application	ns; and
"(III) demonstrate to the	e satisfac-
tion of the Secretary that of	lata from
such records are able to be	exchanged
through the use of applica	tion pro-
gramming interfaces and u	sed in a
manner that allows for exch	ange and
everyday use, as authorized a	under ap-
plicable law, of such records.	
"(ii) Decertification.—U	nder any
program kept or recognized under	· subpara-
graph (A), the Secretary shall en	isure that
beginning January 1, 2019, any	qualified
electronic health records that do n	not satisfy
the certification criteria described	in section
3001(c)(5)(B)(ii) or with respect	to which
the vendor or other entity des	cribed in
clause (i) does not satisfy the req	uirements
under such clause (or is determine	d to be in

1	violation of the terms of the attestation or
2	other requirements under such clause) shall
3	no longer be considered as certified under
4	such program.
5	"(iii) Annual publication.—For
6	2019 and each subsequent year, the Sec-
7	retary shall post on the public Internet
8	website of the Department of Health and
9	Human Services a list of any vendors of or
10	other entities offering qualified electronic
11	health records with respect to which certifi-
12	cation has been withdrawn under clause (ii)
13	during such year.
14	"(iv) Periodic review.—The Sec-
15	retary shall periodically review and confirm
16	that vendors of and other entities offering
17	qualified electronic health records have pub-
18	licly published application programming
19	interfaces and associated documentation as
20	required by clause (i)(II) for purposes of
21	certification and maintaining certification
22	under any program kept or recognized
23	$under\ subparagraph\ (A).$
24	"(v) Pricing information.—For pur-
25	poses of clause $(i)(I)(bb)$, the pricing infor-

1	mation described in this clause, with respect
2	to a vendor of or other entity offering a
3	qualified electronic health record, is the fol-
4	lowing:
5	"(I) Additional types of costs or
6	fees (whether fixed, recurring, trans-
7	action based, or otherwise) imposed by
8	the entity (or any third-party from
9	whom the entity purchases, licenses, or
10	obtains any technology, products, or
11	services in connection with the quali-
12	fied electronic health record) to pur-
13	chase, license, implement, maintain,
14	upgrade, use, or otherwise enable and
15	support the use of capabilities to which
16	such record is to be certified under this
17	section; or in connection with any data
18	generated in the course of using any
19	capability to which the record is to be
20	$so\ certified.$
21	"(II) Limitations, whether by
22	contract or otherwise, on the use of any
23	capability to which the record is to be
24	certified under this section for any
25	purpose within the scope of the record's

1	certification; or in connection with any
2	data generated in the course of using
3	any capability to which the record is
4	to be certified under this section.
5	"(III) Limitations, including
6	technical or practical limitations of
7	technology or its capabilities, that
8	could prevent or impair the successful
9	$implementation, \qquad configuration,$
10	customization, maintenance, support,
11	or use of any capabilities to which the
12	record is to be certified under this sec-
13	tion; or that could prevent or limit the
14	use, exchange, or portability of any
15	data generated in the course of using
16	any capability to which the record is
17	to be so certified.".
18	(2) Additional enforcement provisions
19	UNDER THE PUBLIC HEALTH SERVICE ACT.—Subtitle
20	A of title XXX of the Public Health Service Act (42
21	U.S.C. 300jj-11 et seq.), as amended by subsections
22	(a)(1) and (d), is further amended by adding at the
23	end the following new section:

1 "SEC. 3010A. ENFORCEMENT MECHANISMS.

2	"(a) Inspector General Authority.—The Inspec-
3	tor General of the Department of Health and Human Serv-
4	ices shall have the authority to investigate claims of—
5	"(1) vendors of, or other entities offering, quali-
6	fied electronic health records—
7	"(A) being in violation of an attestation
8	made under section $3001(c)(5)(C)(i)(I)$, with re-
9	spect to the use of such records by a health care
10	provider under a specified meaningful use incen-
11	tive program; and
12	"(B) having engaged in information block-
13	ing (as defined in subsection (f)), unless for a le-
14	gitimate purpose specified by the Secretary, with
15	respect to the use of such records by a health care
16	provider under such a program;
17	"(2) health care providers, with respect to the use
18	of such records under a specified meaningful use in-
19	centive program, having, unless for a legitimate pur-
20	pose specified by the Secretary, engaged in informa-
21	tion blocking (as so defined);
22	"(3) health information system providers de-
23	scribed in subsection (b) having engaged in informa-
24	tion blocking (as so defined), unless for a legitimate
25	purpose specified by the Secretary, with respect to the

1	use of such records under a specified meaningful us	se
2	incentive program; and	

- "(4) vendors of, or other entities offering, health information technology (other than technology described in paragraph (1)), health care providers, with respect to the use of such technology, and health information system providers, with respect to such technology, unless for a legitimate purpose specified by the Secretary, having engaged in information blocking (as so defined).
- "(b) Health Information System Providers.—

 The Inspector General of the Department of Health and

 Human Services shall, in coordination with the Federal

 Trade Commission, ensure that health information system

 providers (such as operators of health information ex
 changes and other systems that facilitate the exchange of

 information) investigate claims of information blocking,

 with respect to the use of such records under a specified

 meaningful use incentive program.

20 "(c) Information Sharing Provisions.—

"(1) IN GENERAL.—The National Coordinator
may serve as a technical consultant to the Inspector
General of the Department of Health and Human
Services and the Federal Trade Commission for purposes of carrying out this section. As such technical

- consultant, the National Coordinator may, notwithstanding any other provision of law, share information related to claims or investigations under subsection (a) or (b) with the Federal Trade Commission for purposes of such investigations.
 - "(2) Protection from disclosure of inforMation.—Any information shared by the National
 Coordinator under paragraph (1) shall not be subject
 to the provisions of section 552 of title 5, United
 States Code (commonly referred to as the Freedom of
 Information Act). Any information acquired pursuant to paragraph (1) shall be held in confidence and
 shall not be disclosed to any person except as may be
 necessary to carry out the purposes of subsection (a).
 - "(3) Non-Application of Paperwork Reduction Act.—Chapter 35 of title 44, United States Code (commonly referred to as the Paperwork Reduction Act of 1995) shall not apply to the National Coordinator or to the Office of the National Coordinator for Health Information Technology with respect to the collection of complaints relating to claims described in subsection (a).
- "(d) Penalty.—Any person or entity determined to have committed an act described in paragraph (1), (2), or (3) of subsection (a), in connection with a specified mean-

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1	ingful use incentive program, shall be subject to a civil mon-
2	etary penalty of not more than \$10,000 for each such act.
3	The provisions of section 1128A (other than subsections (a)
4	and (b)) shall apply to a civil money penalty applied under
5	this subsection in the same manner as they apply to a civil
6	money penalty or proceeding under section $1128A(a)$.
7	"(e) Specified Meaningful Use Incentive Pro-
8	GRAM.—For purposes of this section, the term 'specified
9	meaningful use incentive program' includes the following.
10	"(1) The incentive payments under subsection
11	(o) of section 1848 of the Social Security Act (42
12	U.S.C. 1395w-4) and adjustments under subsection
13	(a) (7) of such section.
14	"(2) The incentive payments under subsection
15	(n) of section 1848 of such Act (42 U.S.C. 1395ww)
16	and adjustments under subsection $(b)(3)(B)$ of such
17	section.
18	"(3) The incentive payments and adjustments
19	made under subsections (l) and (m) of section 1853
20	of such Act (42 U.S.C. 1395w-23).
21	"(4) The incentive payment under paragraph (3)
22	of section 1814(l) of such Act (42 U.S.C. 1395f(l))
23	and adjustment under paragraph (4) of such section.
24	"(5) The shared savings program under section
25	1899 of such Act (42 U.S.C. 1395jjj).

1	"(6) The payments to Medicaid providers de-
2	scribed in section 1903(t) of such Act (42 U.S.C.
3	1396b(t)).
4	"(f) Information Blocking.—
5	"(1) In general.—For purposes of this section
6	and section 3010, the term 'information blocking'
7	means, with respect to the use of qualified electronic
8	health records or other health information technology
9	under a specified meaningful use incentive program,
10	business, technical, and organizational practices, in-
11	cluding practices described in paragraph (2), that—
12	"(A) prevent or materially discourage the
13	exchange of electronic health information;
14	"(B) the actor knows or should know (as de-
15	fined in section $1128A(i)(7)$ of the Social Secu-
16	rity Act) are likely to interfere with the exchange
17	or use of electronic health information; and
18	"(C) do not serve to protect patient safety,
19	maintain the privacy and security of individ-
20	uals' health information or promote competition
21	and consumer welfare.
22	"(2) Practices described.—For purposes of
23	paragraph (1), the practices described in this para-
24	graph are the following:

1	"(A) Contract terms, policies, or other busi-
2	ness or organizational practices that restrict in-
3	dividuals' access to their electronic health infor-
1	mation or restrict the exchange or use of that in-
5	formation for treatment and other permitted
5	purposes.
7	"(B) Charging prices or fees (such as for

- "(B) Charging prices or fees (such as for data exchange, portability, and interfaces) that make exchanging and using electronic health information cost prohibitive.
- "(C) Developing or implementing health information technology in nonstandard ways that are likely to substantially increase the costs, complexity, or burden of sharing electronic health information, especially in cases in which relevant interoperability standards or methods to measure interoperability have been adopted by the Secretary.
- "(D) Developing or implementing health information technology in ways that are likely to lock in users or electronic health information, such as not allowing for the full export of data; lead to fraud, waste, or abuse; or impede innovations and advancements in health information

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1	exchange and health information technology-en-
2	abled care delivery.
3	"(g) Treatment of Vendors With Respect to Pa-
4	TIENT SAFETY ORGANIZATIONS.—In applying part C of
5	title IX—
6	"(1) vendors shall be treated as a provider (as
7	defined in section 921) for purposes of reporting re-
8	quirements under such part, to the extent that such
9	reports are related to attestation requirements under
10	$section \ 3001(c)(5)(C)(i)(I);$
11	"(2) claims of information blocking described in
12	subsection (a) shall be treated as a patient safety ac-
13	tivity under such part for purposes of reporting re-
14	quirements under such part; and
15	"(3) health care providers that are not members
16	of patient safety organizations shall be treated in the
17	same manner as health care providers that are such
18	members for purposes of such reporting requirements
19	with respect to claims of information blocking de-
20	scribed in subsection (a).".
21	(3) ONCHIT.—
22	(A) Portal.—Section 3001(c) of the Public
23	Health Service Act (42 U.S.C. 300jj-11(c)) is
24	amended by adding at the end the following new
25	paragraph:

- "(9) PORTAL.—Not later than January 1, 2019, the National Coordinator shall create a portal to make the information described in paragraph (5)(C)(I)(i)(bb) available to the public in a manner that allows for comparison of price information among health information technology products and that aids in making informed decisions for purchasing such a product."
 - (B) Information blocking.—Not later than 12 months after the date of the enactment of this Act, the National Coordinator of the Office of the National Coordinator of Health Information Technology shall, through rulemaking, implement the provisions of this section, and amendments made by this section, relating to information blocking.
 - (C) HIPAA.—Not later than January 1, 2017, the National Coordinator shall publish guidance to clarify the relationship of the HIPAA privacy and security law, as defined in section 3009(a)(2) of the Public Health Service Act (42 U.S.C. 300jj-19(a)(2)) as such provisions relate to information blocking (as defined in section 3010A(f) of such Act, as added by

1	paragraph (2)), including examples of how such
2	provisions may result in information blocking.
3	(4) Demonstration required for meaning-
4	FUL EHR USE INCENTIVES UNDER MEDICARE.—
5	(A) Incentives for professionals.—
6	(i) In General.—Section
7	1848(o)(2)(C) of the Social Security Act (42
8	U.S.C. 1395 w -4(o)(2)(C)) is amended by
9	adding at the end the following new clause:
10	"(iii) Interoperability.—With re-
11	spect to EHR reporting periods for pay-
12	ment years beginning with 2018, the means
13	described in clause (i) specified by the Sec-
14	retary shall include a demonstration,
15	through means such as an attestation, that
16	the professional has not taken any action
17	described in subsection $(a)(2)$ of section
18	3010A of the Public Health Service Act,
19	with respect to the use of any certified EHR
20	technology.".
21	(ii) Hardship exemption in case of
22	DECERTIFIED EHR.—Subparagraph (B) of
23	section 1848(a)(7) of the Social Security
24	Act (42 U.S.C. 1395w-4(a)(7)) is amended
25	to read as follows:

1	"(B) Significant hardship exception.—
2	"(i) In general.—The Secretary may,
3	on a case-by-case basis, exempt an eligible
4	professional from the application of the
5	payment adjustment under subparagraph
6	(A) if the Secretary determines, subject to
7	annual renewal, that compliance with the
8	requirement for being a meaningful EHR
9	user would result in a significant hardship,
10	such as in the case of an eligible profes-
11	sional who practices in a rural area with-
12	out sufficient Internet access.
13	"(ii) Decertification.—
14	"(I) In General.—The Secretary
15	may, on a case-by-case basis, exempt
16	an eligible professional from the appli-
17	cation of the payment adjustment
18	under subparagraph (A) if the Sec-
19	retary determines that such profes-
20	sional was determined to not be a
21	meaningful EHR user because the
22	qualified electronic health record used
23	by such professional was decertified
24	under section $3001(c)(5)(C)$ of the Pub-

lic Health Service Act. An exemption

1	under the previous sentence may be ap-
2	plied to an eligible professional only,
3	subject to subclause (II), during the
4	first payment year with respect to the
5	first EHR reporting period to which
6	such decertification applies.
7	"(II) Duration.—
8	"(aa) In general.—In no
9	case shall an exemption by reason
10	of this clause be for a period of
11	less than 12 months.
12	"(bb) Extension.—An ex-
13	emption under this clause may be
14	extended for a period of an addi-
15	tional 12 months subject to the
16	limitation described in clause (ii).
17	"(iii) Limitation.—Subject to clause
18	(ii)(II)(aa), in no case may an eligible pro-
19	fessional be granted an exemption under
20	this subparagraph for more than 5 years.".
21	(B) Incentives for hospitals.—
22	(i) In General.—Section 1886(o)(1)
23	of the Social Security Act (42 U.S.C.
24	1395uvv(0)(1)) is amended—

1	(I) in subparagraph (A), by in-
2	serting before the period at the end the
3	following: "and, for performance peri-
4	ods for fiscal year 2018 or a subsequent
5	fiscal year, that provide a demonstra-
6	tion described in subparagraph (D) to
7	the Secretary"; and
8	(II) by adding at the end the fol-
9	lowing new subparagraph:
10	"(D) Demonstration described.—The
11	demonstration described in this subparagraph is
12	a demonstration, through means such as an at-
13	testation, that the hospital has not taken any ac-
14	tion described in subsection $(a)(2)$ of section
15	3010A of the Public Health Service Act, with re-
16	spect to the use of any certified EHR tech-
17	nology.".
18	(ii) Hardship exemption in case of
19	Decertified ehr.—Subclause (II) of sec-
20	tion $1886(b)(3)(B)(ix)$ of the Social Secu-
21	$rity\ Act\ (42\ U.S.C.\ 1395ww(b)(3)(B)(ix))\ is$
22	amended to read as follows:
23	"(II)(aa) The Secretary may, on a case-by-case basis,
24	exempt a subsection (d) hospital from the application of
25	subclause (I) with respect to a fiscal year if the Secretary

- 1 determines, subject to annual renewal, that requiring such
- 2 hospital to be a meaningful EHR user during such fiscal
- 3 year would result in a significant hardship, such as in the
- 4 case of a hospital in a rural area without sufficient Internet
- 5 access.
- 6 "(bb) The Secretary may, on a case-by-case basis, ex-
- 7 empt a subsection (d) hospital from the application of sub-
- 8 clause (I) with respect to a fiscal year if the Secretary deter-
- 9 mines, subject to annual renewal, that such hospital was
- 10 determined to not be a meaningful EHR user because the
- 11 qualified electronic health record used by such hospital was
- 12 decertified under section 3001(c)(5)(C) of the Public Health
- 13 Service Act. An exemption under the previous sentence may
- 14 be applied to a subsection (d) hospital only, subject to items
- 15 (cc) and (dd), during the first payment year with respect
- 16 to the first EHR reporting period to which such decertifica-
- 17 tion applies.
- 18 "(cc) In no case shall an exemption by reason of item
- 19 (bb) be for a period of less than 12 months.
- 20 "(dd) An exemption under item (bb) may be extended
- 21 for a period of an additional 12 months subject to the limi-
- 22 tation described in item (ee).
- 23 "(ee) Subject to item (cc), in no case may a hospital
- 24 be granted an exemption under this subclause for more than
- 25 5 years.".

1 (C) Demonstration required for mean-2 INGFUL EHR USE INCENTIVES UNDER MED-3 ICAID.—Section 1903(t)(2) of the Social Security 4 Act $(42\ U.S.C.\ 1396b(t)(2))$ is amended by add-5 ing at the end the following: "An eligible profes-6 sional shall not qualify as a Medicaid provider 7 under this subsection, with respect to a year be-8 ginning with 2018, unless such provider dem-9 onstrates to the Secretary, through means such as 10 an attestation, that the provider has not taken 11 any action described in subsection (a)(2) of sec-12 tion 3010A of the Public Health Service Act with 13 respect to which the provider knows or should 14 know (as defined in section 1128A(i)(7) of the 15 Social Security Act) about, with respect to the 16 use of any certified EHR technology.".

(f) DEFINITIONS.—

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- (1) Certified ehr technology.—Paragraph
 (1) of section 3000 of the Public Health Service Act
 (42 U.S.C. 300jj) is amended to read as follows:
- "(1) CERTIFIED EHR TECHNOLOGY.—The term
 'certified EHR technology' means a qualified electronic health record that is certified pursuant to section 3001(c)(5) as meeting the certification criteria
 defined in subparagraph (B) of such section that are

1	applicable to the type of record involved (as deter-
2	mined by the Secretary, such as an ambulatory elec-
3	tronic health record for office-based physicians or ar
4	inpatient hospital electronic health record for hos-
5	pitals) including, beginning January 1, 2018, with
6	respect to which the vendor or other entity offering
7	such technology is in compliance with the require
8	ments under section $3001(c)(5)(C)(i)$.".
9	(2) Widespread interoperability.—Section
10	3000 of the Public Health Service Act (42 U.S.C
11	300jj) is amended by adding at the end the following
12	new paragraph:
13	"(15) Widespread interoperability.—The
14	term 'widespread interoperability' means that, on a
15	nationwide basis—
16	"(A) health information technology is inter-
17	operable, in accordance with section 3010; and
18	"(B) such technology is employed by mean
19	ingful EHR users under the specified meaningfu
20	use incentive programs (as defined in section
21	3010A(e)) and by other clinicians and health
22	care providers.".
23	(g) Conforming Amendments.—

1	(1) Voluntary use of standards.—Section
2	3006 of the Public Health Service Act (42 U.S.C.
3	300jj–16) is amended—
4	(A) in subsection (a)(1), by inserting ", in-
5	cluding an interoperability standard adopted
6	under such section" after "section 3004".
7	(B) in subsection (b), by inserting ", in-
8	cluding the interoperability standards adopted
9	under such section" after "section 3004".
10	(2) HIPAA PRIVACY AND SECURITY LAW DEFINI-
11	TION CORRECTION.—Section $3009(a)(2)(A)$ of the
12	Public Health Service Act (42 U.S.C. 300jj-
13	19(a)(2)(A)) is amended by striking "title IV" and
14	inserting "title XIII".
15	(3) Coordination of federal activities.—
16	Section 13111 of the HITECH Act is amended—
17	(A) in subsection (a), by inserting before the
18	period at the end the following: "(and, beginning
19	on January 1, 2018, that are also interoperable
20	under section 3010 of such Act, including by
21	being in compliance with interoperability stand-
22	ards adopted under section 3004 of such Act)";
23	and
24	(B) in subsection (b), by inserting "(and,
25	beginning on January 1, 2018, including an

1	interoperability standard adopted under section
2	3004 of such Act)" before "the President".
3	(4) Application to private entities.—Sec-
4	tion 13112 of the HITECH Act is amended by insert-
5	ing before the period at the end the following: "(and,
6	beginning on January 1, 2018, that are also inter-
7	operable under section 3010 of such Act, including by
8	being in compliance with interoperability standards
9	adopted under section 3004 of such Act)".
10	(5) Coordination with recommendations
11	FOR ACHIEVING WIDESPREAD EHR INTEROPER-
12	ABILITY.—Section 106 of the Medicare Access and
13	CHIP Reauthorization Act of 2015 (Public Law 114-
14	10) is amended by striking subsection (b).".
15	(h) Patient Empowerment.—It is the sense of Con-
16	gress that—
17	(1) patients have the right to the entirety of the
18	health information of such patients, including such
19	information contained in an electronic health record
20	of such patients;
21	(2) such right extends to both structured and
22	unstructured data; and
23	(3) to further facilitate patient ownership over
24	health information of such patient—

1	(A) health care providers should not have
2	the ability to deny a patient's request for access
3	to the entirety of such health information of such
4	patient; and
5	(B) health care providers do not need the
6	consent of their patients to share personal health
7	information of such patients with other covered
8	entities, in compliance with the HIPAA privacy
9	regulations promulgated pursuant to section
10	264(c) of the Health Insurance Portability and
11	Accountability Act of 1996 for the purposes of
12	supporting patient care, except in situations
13	where consent is specifically required under such
14	regulations, such as in cases related to the psy-
15	chiatric records of the patient.
16	Subtitle B—Telehealth
17	SEC. 3021. TELEHEALTH SERVICES UNDER THE MEDICARE
18	PROGRAM.
19	(a) Provision of Information by Centers for
20	Medicare & Medicaid Services.—Not later than 1 year
21	after the date of the enactment of this Act, the Adminis-
22	trator of the Centers for Medicare & Medicaid Services shall
23	provide to the committees of jurisdiction of the House of

24 Representatives and the Senate information on the fol-

25 lowing:

- (1) The populations of Medicare beneficiaries, such as those who are dually eligible for the Medicare program under title XVIII of the Social Security Act (42 U.S.C. 1395 et seg.) and the Medicaid program under title XIX of such Act (42 U.S.C. 1396 et seg.) and those with chronic conditions, whose care may be improved most in terms of quality and efficiency by the expansion, in a manner that meets or exceeds the existing in-person standard of care under the Medi-care program under title XVIII of such Act, of tele-health services under section 1834(m)(4) of such Act $(42\ U.S.C.\ 1395m(m)(4)).$
 - (2) Activities by the Center for Medicare and Medicaid Innovation which examine the use of telehealth services in models, projects, or initiatives funded through section 1115A of the Social Security Act (42 U.S.C. 1315a).
 - (3) The types of high volume procedures codes or diagnoses under such title XVIII which might be suitable to the furnishing of services via telehealth.
 - (4) Barriers that might prevent the expansion of telehealth services under section 1834(m)(4) of the Social Security Act (42 U.S.C. 1395m(m)(4)) beyond such services that are in effect as of the date of the enactment of this Act.

1	(b) Provision of Information by MedPAC.—Not
2	later than 1 year after the date of the enactment of this
3	Act, the Medicare Payment Advisory Commission estab-
4	lished under section 1805 of the Social Security Act (42
5	U.S.C. 1395b-6) shall, using data from the Medicare Ad-
6	vantage program under part C of title XVIII of such Act
7	(42 U.S.C. 1395w-21 et seq.), provide information to the
8	committees of jurisdiction of the House of Representatives
9	and the Senate that identifies—
10	(1) services—
11	(A) for which payment could not be made,
12	as of the date of the enactment of this Act, under
13	the fee-for-service program under parts A and B
14	of such title by reason of any limitation imposed
15	under section 1834(m) of such Act (42 U.S.C.
16	1395m(m)); and
17	(B) that are services that are recommended
18	by the Commission to be included as telehealth
19	services for which payment may be made under
20	the fee-for-service program under parts A and B
21	of such title; and
22	(2) barriers to furnishing telehealth services for
23	which payment may be made under such title XVIII
24	and solutions to address such barriers.

1	(c) Sense of Congress.—It is the sense of Congress
2	that—
3	(1) States should collaborate, through the use of
4	State health board compacts or other mechanisms, to
5	create common licensure requirements services in
6	order to facilitate multistate practices and allow for
7	health care providers to provide such services across
8	State lines;
9	(2) health care providers should be appropriately
10	licensed in the physical location where the patient is
11	receiving services;
12	(3) eligible originating sites should be expanded
13	beyond those originating sites described in section
14	1834(m)(4)(C) of the Social Security Act (42 U.S.C.
15	1395m(m)(4)(C)); and
16	(4) any expansion of telehealth services under the
17	Medicare program should—
18	(A) recognize that telemedicine is the deliv-
19	ery of safe, effective, quality health care services,
20	by a health care provider, using technology as
21	the mode of care delivery;
22	(B) meet or exceed the conditions of cov-
23	erage and payment with respect to the Medicare
24	program under title XVIII unless specifically ad-
25	dress in subsequent statute, of such Act if the

1	service were furnished in person, including
2	standards of care; and
3	(C) involve clinically appropriate means to
4	furnish such services.
5	Subtitle C—Encouraging Con-
6	tinuing Medical Education for
7	Physicians
8	SEC. 3041. EXEMPTING FROM MANUFACTURER TRANS-
9	PARENCY REPORTING CERTAIN TRANSFERS
10	USED FOR EDUCATIONAL PURPOSES.
11	(a) In General.—Section 1128G(e)(10)(B) of the So-
12	cial Security Act (42 U.S.C. 1320a-7h(e)(10)(B)) is
13	amended—
14	(1) in clause (iii), by inserting ", including
15	peer-reviewed journals, journal reprints, journal sup-
16	plements, medical conference reports, and medical
17	textbooks" after "patient use"; and
18	(2) by adding at the end the following new
19	clause:
20	"(xiii) In the case of a covered recipi-
21	ent who is a physician, an indirect pay-
22	ment or transfer of value to the covered re-
23	cipient—
24	"(I) for speaking at, or preparing
25	educational materials for, an edu-

1	cational event for physicians or other
2	health care professionals that does not
3	commercially promote a covered drug,
4	device, biological, or medical supply; or
5	"(II) that serves the sole purpose
6	of providing the covered recipient with
7	medical education, such as by pro-
8	viding the covered recipient with the
9	tuition required to attend an edu-
10	cational event or with materials pro-
11	vided to physicians at an educational
12	event.".
13	(b) Effective Date.—The amendments made by this
14	section shall apply with respect to transfers of value made
15	on or after the date of the enactment of this Act.
16	Subtitle D—Disposable Medical
17	Technologies
18	SEC. 3061. TREATMENT OF CERTAIN ITEMS AND DEVICES.
19	(a) In General.—Section 1834 of the Social Security
20	Act (42 U.S.C. 1395m) is amended by adding at the end
21	the following new subsection:
22	"(r) Payment for Certain Disposable Devices.—
23	"(1) In General.—The Secretary shall make
24	separate payment in the amount established under
25	paragraph (3) to a home health agency for a device

1	described in paragraph (2) when furnished to an in-
2	dividual who receives home health services for which
3	payment is made under section 1895(b).
4	"(2) DEVICE DESCRIBED.—For purposes of
5	paragraph (1), a device described in this paragraph
6	is a disposable device for which, as of January 1,
7	2015, there is—
8	"(A) a Level I Healthcare Common Proce-
9	dure Coding System (HCPCS) code for which
10	the description for a professional service includes
11	the furnishing of such device; and
12	"(B) a separate Level I HCPCS code for a
13	professional service that uses durable medical
14	equipment instead of such device.
15	"(3) Payment amount.—The Secretary shall es-
16	tablish the separate payment amount for such a de-
17	vice such that such amount does not exceed the pay-
18	ment that would be made for the HCPCS code de-
19	scribed in paragraph (2)(A) under section 1833(t)
20	(relating to payment for covered OPD services).".
21	(b) Conforming Amendment.—Section 1861(m)(5)
22	of the Social Security Act (42 U.S.C. $1395x(m)(5)$) is
23	amended by inserting "and devices described in section
24	1834(r)(2)" after "durable medical equipment".

1	(c) Effective Date.—The amendments made by this
2	section shall apply to devices furnished on or after January
3	1, 2017.
4	Subtitle E—Local Coverage
5	Decision Reforms
6	SEC. 3081. IMPROVEMENTS IN THE MEDICARE LOCAL COV-
7	ERAGE DETERMINATION (LCD) PROCESS.
8	(a) In General.—Section 1862(l)(5) of the Social Se-
9	curity Act (42 U.S.C. 1395y(l)(5)) is amended by adding
10	at the end the following new subparagraph:
11	"(D) Local coverage determinations.—
12	The Secretary shall require each medicare ad-
13	ministrative contractor that develops a local cov-
14	erage determination to make available on the
15	website of such contractor and in the coverage
16	database on the Medicare website, at least 45
17	days before the effective date of such determina-
18	tion, the following information:
19	"(i) Such determination in its en-
20	tirety.
21	"(ii) Where and when the proposed de-
22	termination was first made public.
23	"(iii) Hyperlinks to the proposed deter-
24	mination and a response to comments sub-

1	mitted to the contractor with respect to such
2	$proposed\ determination.$
3	"(iv) A summary of evidence that was
4	considered by the contractor during the de-
5	velopment of such determination and a list
6	of the sources of such evidence.
7	"(v) An explanation of the rationale
8	that supports such determination.".
9	(b) Effective Date.—The amendment made by sub-
10	section (a) shall apply with respect to local coverage deter-
11	minations that are proposed or revised on or after the date
12	that is 180 days after the date of the enactment of this Act.
13	Subtitle F-Medicare Pharma-
14	ceutical and Technology Om-
14 15	ceutical and Technology Om- budsman
15	budsman
15 16	budsman SEC. 3101. MEDICARE PHARMACEUTICAL AND TECHNOLOGY
15 16 17	budsman SEC. 3101. MEDICARE PHARMACEUTICAL AND TECHNOLOGY OMBUDSMAN.
15 16 17 18	budsman SEC. 3101. MEDICARE PHARMACEUTICAL AND TECHNOLOGY OMBUDSMAN. Section 1808(c) of the Social Security Act (42 U.S.C.
15 16 17 18 19	budsman SEC. 3101. MEDICARE PHARMACEUTICAL AND TECHNOLOGY OMBUDSMAN. Section 1808(c) of the Social Security Act (42 U.S.C. 1395b-9(c)) is amended by adding at the end the following
15 16 17 18 19 20	budsman SEC. 3101. MEDICARE PHARMACEUTICAL AND TECHNOLOGY OMBUDSMAN. Section 1808(c) of the Social Security Act (42 U.S.C. 1395b-9(c)) is amended by adding at the end the following new paragraph:
15 16 17 18 19 20 21	budsman SEC. 3101. MEDICARE PHARMACEUTICAL AND TECHNOLOGY OMBUDSMAN. Section 1808(c) of the Social Security Act (42 U.S.C. 1395b-9(c)) is amended by adding at the end the following new paragraph: "(4) PHARMACEUTICAL AND TECHNOLOGY OM-
15 16 17 18 19 20 21 22	budsman SEC. 3101. MEDICARE PHARMACEUTICAL AND TECHNOLOGY OMBUDSMAN. Section 1808(c) of the Social Security Act (42 U.S.C. 1395b-9(c)) is amended by adding at the end the following new paragraph: "(4) PHARMACEUTICAL AND TECHNOLOGY OMBUDSMAN.—Not later than 12 months after the date

1	icaid Services who shall receive and respond to com-
2	plaints, grievances, and requests that—
3	"(A) are from entities that manufacture
4	pharmaceutical, biotechnology, medical device, or
5	diagnostic products that are covered or for which
6	coverage is being sought under this title; and
7	"(B) are with respect to coverage, coding, or
8	payment under this title for such products.".
9	Subtitle G—Medicare Site-of-
10	Service Price Transparency
11	SEC. 3121. MEDICARE SITE-OF-SERVICE PRICE TRANS-
12	PARENCY.
13	Section 1834 of the Social Security Act (42 U.S.C.
14	1395m), as amended by section 3061, is further amended
15	by adding at the end the following new subsection:
16	"(s) Site-of-Service Price Transparency.—
17	"(1) In general.—In order to facilitate price
18	transparency with respect to items and services for
19	which payment may be made either to a hospital out-
20	patient department or to an ambulatory surgical cen-
21	ter under this title, the Secretary shall, for 2017 and
22	each year thereafter, make available to the public via
23	a searchable website, with respect to an appropriate
24	number of such items and services—

1	"(A) the estimated payment amount for the
2	item or service under the outpatient department
3	fee schedule under subsection (t) of section 1833
4	and the ambulatory surgical center payment sys-
5	tem under subsection (i) of such section; and
6	"(B) the estimated amount of beneficiary li-
7	ability applicable to the item or service.
8	"(2) Calculation of estimated beneficiary
9	LIABILITY.—For purposes of paragraph (1)(B), the es-
10	timated amount of beneficiary liability, with respect
11	to an item or service, is the amount for such item or
12	service for which an individual who does not have
13	coverage under a medicare supplemental policy cer-
14	tified under section 1882 or any other supplemental
15	insurance coverage is responsible.
16	"(3) Implementation.—In carrying out this
17	subsection, the Secretary—
18	"(A) shall include in the notice described in
19	section 1804(a) a notification of the availability
20	of the estimated amounts made available under
21	paragraph (1); and
22	"(B) may utilize mechanisms in existence
23	on the date of the enactment of this subsection,
24	such as the portion of the website of the Centers
25	for Medicare & Medicaid Services on which in-

1	formation comparing physician performance is
2	posted (commonly referred to as the Physician
3	Compare website), to make available such esti-
4	mated amounts under such paragraph.
5	"(4) Funding.—For purposes of implementing
6	this subsection, the Secretary shall provide for the
7	transfer, from the Supplemental Medical Insurance
8	Trust Fund under section 1841 to the Centers for
9	Medicare & Medicaid Services Program Management
10	Account, of \$6,000,000 for fiscal year 2015, to remain
11	available until expended.".
12	Subtitle H-Medicare Part D Pa-
12	Subtitle II—Medicale Lati D La-
13	tient Safety and Drug Abuse Pre-
13	tient Safety and Drug Abuse Pre-
13 14	tient Safety and Drug Abuse Pre- vention
13 14 15	tient Safety and Drug Abuse Prevention SEC. 3141. PROGRAMS TO PREVENT PRESCRIPTION DRUG
13 14 15 16 17	tient Safety and Drug Abuse Prevention SEC. 3141. PROGRAMS TO PREVENT PRESCRIPTION DRUG ABUSE UNDER MEDICARE PARTS C AND D.
13 14 15 16 17	tient Safety and Drug Abuse Prevention sec. 3141. Programs to prevent prescription drug abuse under medicare parts c and d. (a) Drug Management Program for At-Risk
13 14 15 16 17	tient Safety and Drug Abuse Prevention sec. 3141. Programs to prevent prescription drug abuse under medicare parts c and d. (a) Drug Management Program for At-Risk Beneficiaries.—
13 14 15 16 17 18	tient Safety and Drug Abuse Prevention sec. 3141. Programs to prevent prescription drug Abuse under medicare parts c and d. (a) Drug Management Program for At-Risk Beneficiaries.— (1) In general.—Section 1860D-4(c) of the So-
13 14 15 16 17 18 19 20	tient Safety and Drug Abuse Prevention sec. 3141. Programs to prevent prescription drug Abuse under medicare parts c and d. (a) Drug Management Program for At-Risk Beneficiaries.— (1) In general.—Section 1860D-4(c) of the Social Security Act (42 U.S.C. 1395w-10(c)) is amend-
13 14 15 16 17 18 19 20 21	tient Safety and Drug Abuse Prevention SEC. 3141. PROGRAMS TO PREVENT PRESCRIPTION DRUG ABUSE UNDER MEDICARE PARTS C AND D. (a) Drug Management Program for At-Risk Beneficiaries.— (1) In General.—Section 1860D-4(c) of the Social Security Act (42 U.S.C. 1395w-10(c)) is amended by adding at the end the following:
13 14 15 16 17 18 19 20 21 22	tient Safety and Drug Abuse Prevention SEC. 3141. PROGRAMS TO PREVENT PRESCRIPTION DRUG ABUSE UNDER MEDICARE PARTS C AND D. (a) Drug Management Program for At-Risk Beneficiaries.— (1) In General.—Section 1860D-4(c) of the Social Security Act (42 U.S.C. 1395w-10(c)) is amended by adding at the end the following: "(5) Drug Management Program for At-Risk

1	gram for at-risk beneficiaries under which, sub-
2	ject to subparagraph (B), the PDP sponsor may,
3	in the case of an at-risk beneficiary for prescrip-
4	tion drug abuse who is an enrollee in a prescrip-
5	tion drug plan of such PDP sponsor, limit such
6	beneficiary's access to coverage for frequently
7	abused drugs under such plan to frequently
8	abused drugs that are prescribed for such bene-
9	ficiary by one or more prescribers selected under
10	subparagraph (D), and dispensed for such bene-
11	ficiary by one or more pharmacies selected under
12	such subparagraph.
13	"(B) Requirement for notices.—
14	"(i) In General.—A PDP sponsor
15	may not limit the access of an at-risk bene-
16	ficiary for prescription drug abuse to cov-
17	erage for frequently abused drugs under a
18	prescription drug plan until such sponsor—
19	"(I) provides to the beneficiary an
20	initial notice described in clause (ii)
21	and a second notice described in clause
22	(iii); and
23	"(II) verifies with the providers of
24	the beneficiary that the beneficiary is

1	an at-risk beneficiary for prescription
2	drug abuse.
3	"(ii) Initial notice.—An initial no-
4	tice described in this clause is a notice that
5	provides to the beneficiary—
6	"(I) notice that the PDP sponsor
7	has identified the beneficiary as poten-
8	tially being an at-risk beneficiary for
9	prescription drug abuse;
10	"(II) information describing all
11	State and Federal public health re-
12	sources that are designed to address
13	prescription drug abuse to which the
14	beneficiary has access, including men-
15	tal health services and other counseling
16	services;
17	"(III) notice of, and information
18	about, the right of the beneficiary to
19	appeal such identification under sub-
20	section (h) and the option of an auto-
21	matic escalation to external review;
22	"(IV) a request for the beneficiary
23	to submit to the PDP sponsor pref-
24	erences for which prescribers and phar-
25	macies the beneficiary would prefer the

1	PDP sponsor to select under subpara-
2	graph (D) in the case that the bene-
3	ficiary is identified as an at-risk bene-
4	ficiary for prescription drug abuse as
5	$described \ in \ clause \ (iii)(I);$
6	"(V) an explanation of the mean-
7	ing and consequences of the identifica-
8	tion of the beneficiary as potentially
9	being an at-risk beneficiary for pre-
10	scription drug abuse, including an ex-
11	planation of the drug management
12	program established by the PDP spon-
13	sor pursuant to subparagraph (A);
14	"(VI) clear instructions that ex-
15	plain how the beneficiary can contact
16	the PDP sponsor in order to submit to
17	the PDP sponsor the preferences de-
18	scribed in subclause (IV) and any
19	other communications relating to the
20	drug management program for at-risk
21	beneficiaries established by the PDP
22	sponsor; and
23	"(VII) contact information for
24	other organizations that can provide
25	the beneficiary with assistance regard-

ing such drug management p	program
(similar to the information p	provided
by the Secretary in other stand	dardized
notices provided to part D eliq	gible in-
dividuals enrolled in prescripti	ion drug
plans under this part).	
"(iii) Second notice.—A sec	ond no-
tice described in this clause is a nor	tice that
provides to the beneficiary notice—	
"(I) that the PDP spon	nsor has
identified the beneficiary as ar	ı at-risk
beneficiary for prescription dru	ıg abuse;
"(II) that such beneficiary	j is sub-
ject to the requirements of t	he drug
management program for at-ri	isk bene-
ficiaries established by such PL	P spon-
sor for such plan;	
"(III) of the prescriber	(or pre-
scribers) and pharmacy (or	r phar-
macies) selected for such in	dividual
$under\ subparagraph\ (D);$	
"(IV) of, and information	n about,
the beneficiary's right to appe	eal such
identification under subsection	(h) and

1	the option of an automatic escalation
2	to external review;
3	"(V) that the beneficiary can, in
4	the case that the beneficiary has not
5	previously submitted to the PDP spon-
6	sor preferences for which prescribers
7	and pharmacies the beneficiary would
8	prefer the PDP sponsor select under
9	subparagraph (D), submit such pref-
10	erences to the PDP sponsor; and
11	"(VI) that includes clear instruc-
12	tions that explain how the beneficiary
13	can contact the PDP sponsor.
14	"(iv) Timing of notices.—
15	"(I) In general.—Subject to
16	subclause (II), a second notice de-
17	scribed in clause (iii) shall be provided
18	to the beneficiary on a date that is not
19	less than 60 days after an initial no-
20	tice described in clause (ii) is provided
21	to the beneficiary.
22	"(II) Exception.—In the case
23	that the PDP sponsor, in conjunction
24	with the Secretary, determines that
25	concerns identified through rulemaking

1	by the Secretary regarding the health
2	or safety of the beneficiary or regard-
3	ing significant drug diversion activi-
4	ties require the PDP sponsor to pro-
5	vide a second notice described in clause
6	(iii) to the beneficiary on a date that
7	is earlier than the date described in
8	subclause (I), the PDP sponsor may
9	provide such second notice on such ear-
10	lier date.
11	"(C) At-risk beneficiary for prescrip-
12	TION DRUG ABUSE.—
13	"(i) In general.—For purposes of
14	this paragraph, the term 'at-risk beneficiary
15	for prescription drug abuse' means a part
16	D eligible individual who is not an exempt-
17	ed individual described in clause (ii) and—
18	"(I) who is identified through the
19	use of clinical guidelines developed by
20	the Secretary in consultation with
21	PDP sponsors and other stakeholders
22	described in section $3141(f)(2)(A)$ of
23	the 21st Century Cures Act; or
24	"(II) with respect to whom the
25	PDP sponsor of a prescription drug

1	plan, upon enrolling such individual
2	in such plan, received notice from the
3	Secretary that such individual was
4	identified under this paragraph to be
5	an at-risk beneficiary for prescription
6	drug abuse under the prescription drug
7	plan in which such individual was
8	most recently previously enrolled and
9	such identification has not been termi-
10	nated under subparagraph (F).
11	"(ii) Exempted individual de-
12	SCRIBED.—An exempted individual de-
13	scribed in this clause is an individual
14	who—
15	"(I) receives hospice care under
16	$this\ title;$
17	"(II) is a resident of a long-term
18	care facility, of an intermediate care
19	facility for the mentally retarded, or of
20	another facility for which frequently
21	abused drugs are dispensed for resi-
22	dents through a contract with a single
23	pharmacy; or

1	"(III) the Secretary elects to treat
2	as an exempted individual for purposes
3	of clause (i) .
4	"(D) Selection of prescribers and
5	PHARMACIES.—
6	"(i) In general.—With respect to
7	each at-risk beneficiary for prescription
8	drug abuse enrolled in a prescription drug
9	plan offered by such sponsor, a PDP spon-
10	sor shall, based on the preferences submitted
11	to the PDP sponsor by the beneficiary pur-
12	suant to clauses (ii)(IV) and (iii)(V) of sub-
13	paragraph (B), select—
14	"(I) one or more individuals who
15	are authorized to prescribe frequently
16	abused drugs (referred to in this para-
17	graph as 'prescribers') who may write
18	prescriptions for such drugs for such
19	beneficiary; and
20	"(II) one or more pharmacies that
21	may dispense such drugs to such bene-
22	ficiary.
23	"(ii) Reasonable access.—In mak-
24	ing the selections under this subpara-
25	graph—

1	"(I) a PDP sponsor shall ensure
2	that the beneficiary continues to have
3	reasonable access to frequently abused
4	drugs (as defined in subparagraph
5	(G)), taking into account geographic
6	location, beneficiary preference, impact
7	on costsharing, and reasonable travel
8	time; and
9	"(II) a PDP sponsor shall ensure
10	such access (including access to pre-
11	scribers and pharmacies with respect
12	to frequently abused drugs) in the case
13	of individuals with multiple residences
14	and in the case of natural disasters
15	and similar emergency situations.
16	"(iii) Beneficiary preferences.—
17	"(I) In general.—If an at-risk
18	beneficiary for prescription drug abuse
19	submits preferences for which in-net-
20	work prescribers and pharmacies the
21	beneficiary would prefer the PDP
22	sponsor select in response to a notice
23	under subparagraph (B), the PDP
24	sponsor shall—

"(aa) review such	pref-
erences;	
"(bb) select or change	the se-
lection of prescribers and	phar-
macies for the beneficiary	based
on such preferences; and	
"(cc) inform the benef	ficiary
of such selection or change	of se-
lection.	
"(II) Exception.—In the	e case
that the PDP sponsor determine	es that
a change to the selection of pre-	scriber
or pharmacy under item (bb)	by the
PDP sponsor is contributing or	would
contribute to prescription drug	abuse
or drug diversion by the benef	ïciary,
the PDP sponsor may change the	e selec-
tion of prescriber or pharmacy j	for the
beneficiary without regard to the	e pref-
erences of the beneficiary descri	bed in
subclause (I).	
"(iv) Confirmation.—Before se	lecting
a prescriber (or prescribers) or pha	rmacy
(or pharmacies) under this subparagr	aph, a
PDP sponsor must request and receiv	ve con-

1	firmation from such a prescriber or phar-
2	macy acknowledging and accepting that the
3	beneficiary involved is in the drug manage-
4	ment program for at-risk beneficiaries.
5	"(E) TERMINATIONS AND APPEALS.—The
6	identification of an individual as an at-risk ben-
7	eficiary for prescription drug abuse under this
8	paragraph, a coverage determination made
9	under a drug management program for at-risk
10	beneficiaries, and the selection of prescriber or
11	pharmacy under subparagraph (D) with respect
12	to such individual shall be subject to reconsider-
13	ation and appeal under subsection (h) and the
14	option of an automatic escalation to external re-
15	view to the extent provided by the Secretary.
16	"(F) TERMINATION OF IDENTIFICATION.—
17	"(i) In General.—The Secretary shall
18	develop standards for the termination of
19	identification of an individual as an at-risk
20	beneficiary for prescription drug abuse
21	under this paragraph. Under such stand-
22	ards such identification shall terminate as
23	of the earlier of—
24	"(I) the date the individual dem-
25	onstrates that the individual is no

1	longer likely, in the absence of the re-
2	strictions under this paragraph, to be
3	an at-risk beneficiary for prescription
4	drug abuse described in subparagraph
5	(C)(i); and
6	"(II) the end of such maximum
7	period of identification as the Sec-
8	retary may specify.
9	"(ii) Rule of construction.—Noth-
10	ing in clause (i) shall be construed as pre-
11	venting a plan from identifying an indi-
12	vidual as an at-risk beneficiary for pre-
13	scription drug abuse under subparagraph
14	(C)(i) after such termination on the basis of
15	additional information on drug use occur-
16	ring after the date of notice of such termi-
17	nation.
18	"(G) Frequently abused drug.—For
19	purposes of this subsection, the term 'frequently
20	abused drug' means a drug that is a controlled
21	substance that the Secretary determines to be fre-
22	quently abused or diverted.
23	"(H) Data disclosure.—In the case of an
24	at-risk beneficiary for prescription drug abuse
25	whose access to coverage for frequently abused

1	drugs under a prescription drug plan has been
2	limited by a PDP sponsor under this paragraph,
3	such PDP sponsor shall disclose data, including
4	any necessary individually identifiable health
5	information, in a form and manner specified by
6	the Secretary, about the decision to impose such
7	limitations and the limitations imposed by the
8	sponsor under this part.
9	"(I) Education.—The Secretary shall pro-
10	vide education to enrollees in prescription drug
11	plans of PDP sponsors and providers regarding
12	the drug management program for at-risk bene-
13	ficiaries described in this paragraph, including
14	education—
15	"(i) provided by medicare administra-
16	tive contractors through the improper pay-
17	ment outreach and education program de-
18	scribed in section 1874A(h); and
19	"(ii) through current education efforts
20	(such as State health insurance assistance
21	$programs\ described\ in\ subsection\ (a)(1)(A)$
22	of section 119 of the Medicare Improvements
23	for Patients and Providers Act of 2008 (42
24	U.S.C. 1395b-3 note)) and materials di-
25	rected toward such enrollees.

1	"(J) APPLICATION UNDER MA-PD PLANS.—
2	Pursuant to section 1860D—21(c)(1), the provi-
3	sions of this paragraph apply under part D to
4	MA organizations offering MA-PD plans to MA
5	eligible individuals in the same manner as such
6	provisions apply under this part to a PDP spon-
7	sor offering a prescription drug plan to a part
8	D eligible individual.".
9	(2) Information for consumers.—Section
10	1860D-4(a)(1)(B) of the Social Security Act (42)
11	U.S.C. $1395w-104(a)(1)(B)$) is amended by adding at
12	the end the following:
13	"(v) The drug management program
14	for at-risk beneficiaries under subsection
15	(c)(5).".
16	(b) Utilization Management Programs.—Section
17	1860D-4(c) of the Social Security Act (42 U.S.C. 1395w-
18	104(c)), as amended by subsection (a)(1), is further amend-
19	ed—
20	(1) in paragraph (1), by inserting after subpara-
21	graph (D) the following new subparagraph:
22	"(E) A utilization management tool to pre-
23	vent drug abuse (as described in paragraph
24	(6)(A)).": and

1	(2) by adding at the end the following new para-
2	graph:
3	"(6) Utilization management tool to pre-
4	VENT DRUG ABUSE.—
5	"(A) In general.—A tool described in this
6	paragraph is any of the following:
7	"(i) A utilization tool designed to pre-
8	vent the abuse of frequently abused drugs by
9	individuals and to prevent the diversion of
10	such drugs at pharmacies.
11	"(ii) Retrospective utilization review to
12	identify—
13	"(I) individuals that receive fre-
14	quently abused drugs at a frequency or
15	in amounts that are not clinically ap-
16	propriate; and
17	"(II) providers of services or sup-
18	pliers that may facilitate the abuse or
19	diversion of frequently abused drugs by
20	beneficiaries.
21	"(iii) Consultation with the contractor
22	described in subparagraph (B) to verify if
23	an individual enrolling in a prescription
24	drug plan offered by a PDP sponsor has
25	been previously identified by another PDP

1	sponsor as an individual described in clause
2	(ii)(I).
3	"(B) Reporting.—A PDP sponsor offering
4	a prescription drug plan (and an MA organiza-
5	tion offering an MA-PD plan) in a State shall
6	submit to the Secretary and the Medicare drug
7	integrity contractor with which the Secretary
8	has entered into a contract under section 1893
9	with respect to such State a report, on a monthly
10	basis, containing information on—
11	"(i) any provider of services or sup-
12	plier described in subparagraph (A)(ii)(II)
13	that is identified by such plan sponsor (or
14	organization) during the 30-day period be-
15	fore such report is submitted; and
16	"(ii) the name and prescription
17	records of individuals described in para-
18	$graph\ (5)(C)$.".
19	(c) Expanding Activities of Medicare Drug In-
20	TEGRITY CONTRACTORS (MEDICs).—
21	(1) In General.—Section 1893 of the Social Se-
22	curity Act (42 U.S.C. 1395ddd) is amended by add-
23	ing at the end the following new subsection:
24	"(j) Expanding Activities of Medicare Drug In-
25	TEGRITY CONTRACTORS (MEDICS).—

1	"(1) Access to information.—Under contracts
2	entered into under this section with Medicare drug
3	integrity contractors (including any successor entity
4	to a Medicare drug integrity contractor), the Sec-
5	retary shall authorize such contractors to directly ac-
6	cept prescription and necessary medical records from
7	entities such as pharmacies, prescription drug plans,
8	MA-PD plans, and physicians with respect to an in-
9	dividual in order for such contractors to provide in-
10	formation relevant to the determination of whether
11	such individual is an at-risk beneficiary for prescrip-
12	tion drug abuse, as defined in section 1860D-
13	4(c)(5)(C).
14	"(2) Requirement for acknowledgment of
15	REFERRALS.—If a PDP sponsor or MA organization
16	refers information to a contractor described in para-
17	graph (1) in order for such contractor to assist in the
18	determination described in such paragraph, the con-
19	tractor shall—
20	"(A) acknowledge to the sponsor or organi-
21	zation receipt of the referral; and
22	"(B) in the case that any PDP sponsor or
23	MA organization contacts the contractor request-
24	ing to know the determination by the contractor
25	of whether or not an individual has been deter-

1	mined to be an individual described such para-
2	graph, shall inform such sponsor or organization
3	of such determination on a date that is not later
4	than 15 days after the date on which the sponsor
5	or organization contacts the contractor.
6	"(3) Making data available to other enti-
7	TIES.—
8	"(A) In general.—For purposes of car-
9	rying out this subsection, subject to subpara-
10	graph (B), the Secretary shall authorize MED-
11	ICs to respond to requests for information from
12	PDP sponsors and MA organizations, State pre-
13	scription drug monitoring programs, and other
14	entities delegated by such sponsors or organiza-
15	tions using available programs and systems in
16	the effort to prevent fraud, waste, and abuse.
17	"(B) HIPAA COMPLIANT INFORMATION
18	ONLY.—Information may only be disclosed by a
19	MEDIC under subparagraph (A) if the disclo-
20	sure of such information is permitted under the
21	Federal regulations (concerning the privacy of
22	individually identifiable health information)
23	promulgated under section 264(c) of the Health
24	Insurance Portability and Accountability Act of

1996 (42 U.S.C. 1320d–2 note).".

25

1	(2) OIG STUDY AND REPORT ON EFFECTIVENESS
2	OF MEDICS.—
3	(A) Study.—The Inspector General of the
4	Department of Health and Human Services shall
5	conduct a study on the effectiveness of Medicare
6	drug integrity contractors with which the Sec-
7	retary of Health and Human Services has en-
8	tered into a contract under section 1893 of the
9	Social Security Act (42 U.S.C. 1395ddd) in
10	identifying, combating, and preventing fraud
11	under the Medicare program, including under
12	the authority provided under section 1893(j) of
13	the Social Security Act, added by paragraph (1).
14	(B) Report.—Not later than 1 year after
15	the date of the enactment of this Act, the Inspec-
16	tor General shall submit to Congress a report on
17	the study conducted under subparagraph (A).
18	Such report shall include such recommendations
19	for improvements in the effectiveness of such con-
20	tractors as the Inspector General determines ap-
21	propriate.
22	(d) Treatment of Certain Complaints for Pur-
23	Poses of Quality or Performance Assessment.—Sec-
24	tion 1860D-42 of the Social Security Act (42 U.S.C.

- 1 1395w-152) is amended by adding at the end the following
- 2 new subsection:
- 3 "(d) Treatment of Certain Complaints for Pur-
- 4 Poses of Quality or Performance Assessment.—In
- 5 conducting a quality or performance assessment of a PDP
- 6 sponsor, the Secretary shall develop or utilize existing
- 7 screening methods for reviewing and considering com-
- 8 plaints that are received from enrollees in a prescription
- 9 drug plan offered by such PDP sponsor and that are com-
- 10 plaints regarding the lack of access by the individual to
- 11 prescription drugs due to a drug management program for
- 12 at-risk beneficiaries.".
- 13 (e) Sense of Congress Regarding Use of Tech-
- 14 NOLOGY TOOLS TO COMBAT FRAUD.—It is the sense of Con-
- 15 gress that MA organizations and PDP sponsors should con-
- 16 sider using e-prescribing and other health information tech-
- 17 nology tools to support combating fraud under MA-PD
- 18 plans and prescription drug plans under parts C and D
- 19 of the Medicare program.
- 20 (f) Effective Date.—
- 21 (1) In General.—The amendments made by
- 22 this section shall apply to prescription drug plans
- 23 (and MA-PD plans) for plan years beginning more
- 24 than 1 year after the date of the enactment of this
- 25 *Act*.

1	(2) Stakeholder meetings prior to effec-
2	TIVE DATE.—
3	(A) In general.—Not later than January
4	1, 2016, the Secretary of Health and Human
5	Services shall convene stakeholders, including in-
6	dividuals entitled to benefits under part A of
7	title XVIII of the Social Security Act or enrolled
8	under part B of such title of such Act, advocacy
9	groups representing such individuals, physicians,
10	pharmacists, and other clinicians, retail phar-
11	macies, plan sponsors, entities delegated by plan
12	sponsors, and biopharmaceutical manufacturers
13	for input regarding the topics described in sub-
14	paragraph (B).
15	(B) Topics described.—The topics de-
16	scribed in this subparagraph are the topics of—
17	(i) the impact on cost-sharing and en-
18	suring accessibility to prescription drugs for
19	enrollees in prescription drug plans of PDP
20	sponsors, and enrollees in MA-PD plans,
21	who are at-risk beneficiaries for prescrip-
22	tion drug abuse (as defined in subpara-
23	graph (C) of paragraph (5) of section
24	1860D-4(c) of the Social Security Act (42
25	$U.S.C.\ 1395w-104(c));$

1	(ii) the use of an expedited appeals
2	process under which such an enrollee may
3	appeal an identification of such enrollee as
4	an at-risk beneficiary for prescription drug
5	abuse under such paragraph (similar to the
6	processes established under the Medicare Ad-
7	vantage program under part C of title
8	XVIII of the Social Security Act that allow
9	an automatic escalation to external review
10	of claims submitted under such part);
11	(iii) the types of enrollees that should
12	be treated as exempted individuals, as de-
13	scribed in subparagraph (C)(ii) of such
14	paragraph;
15	(iv) the manner in which terms and
16	definitions in such paragraph should be ap-
17	plied, such as the use of clinical appro-
18	priateness in determining whether an en-
19	rollee is an at-risk beneficiary for prescrip-
20	tion drug abuse as defined in subparagraph
21	(C) of such paragraph;
22	(v) the information to be included in
23	the notices described in subparagraph (B) of
24	such paragraph and the standardization of
25	such notices; and

1	(vi) with respect to a PDP sponsor (or
2	Medicare Advantage organization) that es-
3	tablishes a drug management program for
4	at-risk beneficiaries under such paragraph,
5	the responsibilities of such PDP sponsor (or
6	organization) with respect to the implemen-
7	tation of such program.
8	(g) Rulemaking.—The Secretary of Health and
9	Human Services shall promulgate regulations based on the
10	input gathered pursuant to subsection $(f)(2)(A)$.
11	TITLE IV—MEDICAID, MEDICARE,
12	AND OTHER REFORMS
13	Subtitle A—Medicaid and Medicare
14	Reforms
15	SEC. 4001. LIMITING FEDERAL MEDICAID REIMBURSEMENT
16	TO STATES FOR DURABLE MEDICAL EQUIP-
17	MENT (DME) TO MEDICARE PAYMENT RATES.
18	(a) Medicaid Reimbursement.—
19	(1) In General.—Section 1903(i) of the Social
20	Security Act (42 U.S.C. 1396b(i)) is amended—
21	(A) in paragraph (25), by striking "or" at
22	$the\ end;$
23	(B) in paragraph (26), by striking the pe-
24	riod at the end and inserting "; or"; and

1	(C) by inserting after paragraph (26) the
2	following new paragraph:
3	"(27) with respect to any amounts expended by
4	the State on the basis of a fee schedule for items de-
5	scribed in section 1861(n), as determined in the ag-
6	gregate with respect to each class of such items as de-
7	fined by the Secretary, in excess of the aggregate
8	amount, if any, that would be paid for such items
9	within such class on a fee-for-service basis under the
10	program under part B of title XVIII, including, as
11	applicable, under a competitive acquisition program
12	under section 1847 in an area of the State.".
13	(2) Effective date.—The amendments made
14	by this subsection shall be effective with respect to
15	payments for items furnished on or after January 1,
16	2020.
17	(b) Medicare Ombudsman.—Section 1808(c) of the
18	Social Security Act (42 U.S.C. 1395b(c)), as amended by
19	section 3101, is further amended by adding at the end the
20	following new paragraph:
21	"(5) Monitoring dme reimbursement under
22	MEDICAID.—The ombudsmen under each of para-
23	graphs (1) and (4) shall evaluate the impact of the
24	competitive acquisition program under section 1847,

1	including as applied under section $1903(i)(27)$, on
2	beneficiary health status and health outcomes.".
3	SEC. 4002. MEDICARE PAYMENT INCENTIVE FOR THE TRAN-
4	SITION FROM TRADITIONAL X-RAY IMAGING
5	TO DIGITAL RADIOGRAPHY AND OTHER MEDI-
6	CARE IMAGING PAYMENT PROVISION.
7	(a) Physician Fee Schedule.—
8	(1) Payment incentive for transition.—
9	(A) In General.—Section 1848(b) of the
10	Social Security Act (42 U.S.C. 1395w-4(b)) is
11	amended by adding at the end the following new
12	paragraph:
13	"(9) Special rule to incentivize transition
14	FROM TRADITIONAL X-RAY IMAGING TO DIGITAL RADI-
15	OGRAPHY.—
16	"(A) Limitation on payment for film x-
17	RAY IMAGING SERVICES.—In the case of imaging
18	services that are X rays taken using film and
19	that are furnished during 2017 or a subsequent
20	year, the payment amount for the technical com-
21	ponent (including the technical component por-
22	tion of a global fee) of such services that would
23	otherwise be determined under this section (with-
24	out application of this paragraph and before ap-
25	plication of any other adjustment under this sec-

1	tion) for such year shall be reduced by 20 per-
2	cent.
3	"(B) Phased-in limitation on payment
4	FOR COMPUTED RADIOGRAPHY IMAGING SERV-
5	ICES.—In the case of imaging services that are
6	X rays taken using computed radiography tech-
7	nology—
8	"(i) in the case of such services fur-
9	nished during 2018, 2019, 2020, 2021, or
10	2022 the payment amount for the technical
11	component (including the technical compo-
12	nent portion of a global fee) of such services
13	that would otherwise be determined under
14	this section (without application of this
15	paragraph and before application of any
16	other adjustment under this section) for
17	such year shall be reduced by 7 percent; and
18	"(ii) in the case of such services fur-
19	nished during 2023 or a subsequent year,
20	the payment amount for the technical com-
21	ponent (including the technical component
22	portion of a global fee) of such services that
23	would otherwise be determined under this
24	section (without application of this para-
25	graph and before application of any other

1	adjustment under this section) for such year
2	shall be reduced by 10 percent.
3	"(C) Computed Radiography tech-
4	NOLOGY DEFINED.—For purposes of this para-
5	graph, the term 'computed radiography tech-
6	nology' means cassette-based imaging which uti-
7	lizes an imaging plate to create the image in-
8	volved.
9	"(D) Implementation.—In order to imple-
10	ment this paragraph, the Secretary shall adopt
11	appropriate mechanisms which may include use
12	of modifiers.".
13	(B) Exemption from budget neu-
14	TRALITY.—Section $1848(c)(2)(B)(v)$ of the Social
15	Security Act (42 U.S.C. $1395w-4(c)(2)(B)(v)$) is
16	amended by adding at the end the following new
17	subclause:
18	"(X) Reduced expenditures
19	ATTRIBUTABLE TO INCENTIVES TO
20	TRANSITION TO DIGITAL RADIOG-
21	RAPHY.—Effective for fee schedules es-
22	tablished beginning with 2017, reduced
23	expenditures attributable to subpara-
24	graph (A) of subsection (b)(9) and ef-
25	fective for fee schedules established be-

1	ginning with 2018, reduced expendi-
2	tures attributable to subparagraph (B)
3	of such subsection.".
4	(2) Elimination of application of multiple
5	PROCEDURE PAYMENT REDUCTION.—Section
6	1848(b)(4) of the Social Security Act (42 U.S.C.
7	1395w-4(b)(4)) is amended by adding at the end the
8	following new subparagraph:
9	"(E) Elimination of Application of
10	MULTIPLE PROCEDURE PAYMENT REDUCTION.—
11	"(i) In general.—Not later than Jan-
12	uary 1, 2016, the Secretary shall not apply
13	a multiple procedure payment reduction
14	policy to the professional component of im-
15	aging services furnished in any subsequent
16	year that is prior to a year in which the
17	Secretary conducts and publishes, as part of
18	the Medicare Physician Fee Schedule Pro-
19	posed Rule for a year, the empirical anal-
20	ysis described in clause (ii).
21	"(ii) Empirical analysis de-
22	SCRIBED.—The empirical analysis described
23	in this clause is an analysis of the Re-
24	source-Based Relative Value Scale (com-
25	monly known as the 'RBRVS') Data Man-

1	ager information that is used to determine
2	what, if any, efficiencies exist within the
3	professional component of imaging services
4	when two or more studies are performed on
5	the same patient on the same day. Such em-
6	pirical analysis shall include—
7	"(I) work sheets and other infor-
8	mation detailing which physician work
9	activities performed given the typical
10	vignettes were assigned reduction per-
11	centages of 0, 25, 50, 75 and 100 per-
12	cent;
13	"(II) a discussion of the clinical
14	aspects that informed the assignment of
15	the reduction percentages described in
16	subclause (I);
17	"(III) an explanation of how the
18	percentage reductions for pre-, intra-,
19	and post-service work were determined
20	and calculated; and
21	"(IV) a demonstration that the
22	Centers for Medicare & Medicaid Serv-
23	ices has consulted with practicing ra-
24	diologists to gain knowledge of how ra-
25	diologists interpret studies of multiple

1	body parts on the same individual on
2	the same day.".
3	(b) Payment Incentive for Transition Under
4	Hospital Outpatient Prospective Payment Sys-
5	TEM.—Section 1833(t)(16) of the Social Security Act (42
6	$U.S.C.\ 1395(t)(16))$ is amended by adding at the end the
7	following new subparagraph:
8	"(F) Payment incentive for the transi-
9	TION FROM TRADITIONAL X-RAY IMAGING TO DIG-
10	ITAL RADIOGRAPHY.—Notwithstanding the pre-
11	vious provisions of this subsection:
12	"(i) Limitation on payment for
13	FILM X-RAY IMAGING SERVICES.—In the
14	case of imaging services that are X rays
15	taken using film and that are furnished
16	during 2017 or a subsequent year, the pay-
17	ment amount for the technical component
18	(including the technical component portion
19	of a global fee) of such services that would
20	otherwise be determined under this section
21	(without application of this paragraph and
22	before application of any other adjustment
23	under this subsection) for such year shall be
24	reduced by 20 percent.

1 "(ii) Phased-in limitation on pay
MENT FOR COMPUTED RADIOGRAPHY IMAG
3 ING SERVICES.—In the case of imaging
4 services that are X rays taken using com
5 puted radiography technology (as defined in
section 1848(b)(9)(C))—
"(I) in the case of such service.
furnished during 2018, 2019, 2020
2021, or 2022 the payment amount for
the technical component (including the
1 technical component portion of a globa
fee) of such services that would other
3 wise be determined under this section
4 (without application of this paragraph
and before application of any other ad
justment under this subsection) for
such year shall be reduced by 7 per
S cent; and
"(II) in the case of such service.
furnished during 2023 or a subsequen
1 year, the payment amount for the tech
2 nical component (including the tech
nical component portion of a globa
fee) of such services that would other
5 wise be determined under this section

1	(without application of this paragraph
2	and before application of any other ad-
3	justment under this subsection) for
4	such year shall be reduced by 10 per-
5	cent.
6	"(iii) Application without regard
7	TO BUDGET NEUTRALITY.—The reductions
8	made under this paragraph—
9	"(I) shall not be considered an ad-
10	justment under paragraph $(2)(E)$; and
11	"(II) shall not be implemented in
12	a budget neutral manner.".
13	SEC. 4003. IMPLEMENTATION OF OFFICE OF INSPECTOR
14	GENERAL RECOMMENDATION TO DELAY CER-
15	TAIN MEDICARE PRESCRIPTION DRUG PLAN
16	PREPAYMENTS.
17	Section 1860D-15(d) of the Social Security Act (42
18	U.S.C. 1395w-115(d)) is amended by adding at the end the
19	following:
20	"(5) Timing of payments.—With respect to
21	monthly reinsurance payment amounts under this
22	section to a PDP sponsor for months in a year (be-
23	ginning with 2020), such payment amounts for a
24	month shall be made on the first business day occur-
25	ring on or after the following date for that month:

1	"(A) For the month of January, January
2	2nd.
3	"(B) For the month of February, February
4	5th.
5	"(C) For the month of March, March 10th.
6	"(D) For the month of April, April 15th.
7	"(E) For the month of May, May 20th.
8	"(F) For the month of June, June 25th.
9	"(G) For the month of July and each suc-
10	ceeding month (other than December) in a year,
11	the first day of the next month.
12	"(H) For the month of December, December
13	24th.".
14	Subtitle B—Cures Innovation Fund
15	SEC. 4041. CURES INNOVATION FUND.
16	(a) Establishment.—There is hereby established in
17	the Treasury of the United States a fund to be known as
18	the Cures Innovation Fund (in this section referred to as
19	the "Fund").
20	(b) APPROPRIATIONS.—There is hereby appropriated
21	to the Fund, out of any funds in the Treasury not otherwise
22	appropriated, \$110,000,000 for each of fiscal years 2016
23	through 2020.

1	(c) Expenditures.—Amounts in the Fund shall be
2	available, as provided by appropriation Acts, for making
3	expenditures for carrying out the following:
4	(1) Section 229A of the Public Health Service
5	Act, as added by section 1123 (relating to data on
6	natural history of diseases).
7	(2) Part E of title II of the Public Health Serv-
8	ice Act, as added by section 1141 (relating to Council
9	for 21st Century Cures).
10	(3) Section 2001 and the amendments made by
11	such section (relating to development and use of pa-
12	tient experience data to enhance structured risk-ben-
13	efit assessment framework).
14	(4) Section 2021 and the amendments made by
15	such section (relating to qualification of drug develop-
16	ment tools).
17	(5) Section 2062 and the amendments made by
18	such section (relating to utilizing evidence from clin-
19	ical experience).
20	(6) Section 2161 (relating to grants for studying
21	the process of continuous drug manufacturing).
22	(d) Supplement, Not Supplant; Prohibition
23	AGAINST TRANSFER.—Funds appropriated by subsection
24	(b)—

1	(1) shall be used to supplement, not supplant,
2	amounts otherwise made available to the National In-
3	stitutes of Health and the Food and Drug Adminis-
4	tration; and
5	(2) notwithstanding any transfer authority in
6	any appropriation Act, shall not be used for any pur-
7	pose other than the expenditures listed in subsection
8	(c).
9	Subtitle C—Other Reforms
10	SEC. 4061. SPR DRAWDOWN.
11	(a) Drawdown and Sale.—Notwithstanding section
12	161 of the Energy Policy and Conservation Act (42 U.S.C.
13	6241), the Secretary of Energy shall draw down and self
14	8,000,000 barrels of crude oil from the Strategic Petroleum
15	Reserve during each of the fiscal years 2018 through 2025,
16	except as provided in subsection (b). Amounts received for
17	a sale under this subsection shall be deposited in the general
18	fund of the Treasury during the fiscal year in which the
19	sale occurs.
20	(b) Emergency Protection.—The Secretary shall
21	not draw down and sell crude oil under this section in
22	amounts that would result in a Strategic Petroleum Reserve
23	that contains an inventory of petroleum products rep-

24 resenting less than 90 days of emergency reserves, based on

- 1 the average daily level of net imports of crude oil and petro-
- 2 leum products in the previous calendar year.
- 3 (c) Proceeds.—Proceeds from a sale under this sec-
- 4 tion shall be deposited into the general fund of the Treasury
- 5 of the United States.

6 Subtitle D—Miscellaneous

- 7 SEC. 4081. LYME DISEASE AND OTHER TICK-BORNE DIS-
- 8 EASES.
- 9 (a) In General.—Title III of the Public Health Serv-
- 10 ice Act (42 U.S.C. 241 et seq.) is amended by adding at
- 11 the end the following new part:
- 12 "PART W—LYME DISEASE AND OTHER TICK-
- 13 BORNE DISEASES
- 14 "SEC. 39900. RESEARCH.
- 15 "(a) In General.—The Secretary shall conduct or
- 16 support epidemiological, basic, translational, and clinical
- 17 research regarding Lyme disease and other tick-borne dis-
- 18 eases.
- 19 "(b) Biennial Reports.—The Secretary shall ensure
- 20 that each biennial report under section 403 includes infor-
- 21 mation on actions undertaken by the National Institutes
- 22 of Health to carry out subsection (a) with respect to Lyme
- 23 disease and other tick-borne diseases, including an assess-
- 24 ment of the progress made in improving the outcomes of
- 25 Lyme disease and such other tick-borne diseases.

"SEC. 39900-1. WORKING GROUP.

2	"(a) Establishment.—The Secretary shall establish
3	a permanent working group, to be known as the Interagency
4	Lyme and Tick-Borne Disease Working Group (in this sec-
5	tion and section 39900-2 referred to as the Working
6	Group'), to review all efforts within the Department of
7	Health and Human Services concerning Lyme disease and
8	other tick-borne diseases to ensure interagency coordination,
9	minimize overlap, and examine research priorities.
10	"(b) Responsibilities.—The Working Group shall—
11	"(1) not later than 24 months after the date of
12	enactment of this part, and every 24 months there-
13	after, develop or update a summary of—
14	"(A) ongoing Lyme disease and other tick-
15	borne disease research related to causes, preven-
16	tion, treatment, surveillance, diagnosis,
17	diagnostics, duration of illness, intervention, and
18	access to services and supports for individuals
19	with Lyme disease or other tick-borne diseases;
20	"(B) advances made pursuant to such re-
21	search;
22	"(C) the engagement of the Department of
23	Health and Human Services with persons that
24	participate at the public meetings required by
25	paragraph (5); and

1	"(D) the comments received by the Working
2	Group at such public meetings and the Sec-
3	retary's response to such comments;
4	"(2) ensure that a broad spectrum of scientific
5	viewpoints is represented in each such summary;
6	"(3) monitor Federal activities with respect to
7	Lyme disease and other tick-borne diseases;
8	"(4) make recommendations to the Secretary re-
9	garding any appropriate changes to such activities;
10	and
11	"(5) ensure public input by holding annual pub-
12	lic meetings that address scientific advances, research
13	questions, surveillance activities, and emerging
14	strains in species of pathogenic organisms.
15	"(c) Membership.—
16	"(1) In general.—The Working Group shall be
17	composed of a total of 14 members as follows:
18	"(A) FEDERAL MEMBERS.—Seven Federal
19	members, consisting of one or more representa-
20	tives of each of—
21	"(i) the Office of the Assistant Sec-
22	retary for Health;
23	"(ii) the Food and Drug Administra-
24	tion;

1	"(iii) the Centers for Disease Control
2	and Prevention;
3	"(iv) the National Institutes of Health;
4	and
5	"(v) such other agencies and offices of
6	the Department of Health and Human
7	Services as the Secretary determines appro-
8	priate.
9	"(B) Non-federal public members.—
10	Seven non-Federal public members, consisting of
11	representatives of the following categories:
12	"(i) Physicians and other medical pro-
13	viders with experience in diagnosing and
14	treating Lyme disease and other tick-borne
15	diseases.
16	"(ii) Scientists or researchers with ex-
17	pertise.
18	"(iii) Patients and their family mem-
19	bers.
20	"(iv) Nonprofit organizations that ad-
21	vocate for patients with respect to Lyme
22	disease and other tick-borne diseases.
23	"(v) Other individuals whose expertise
24	is determined by the Secretary to be bene-

1	ficial to the functioning of the Working
2	Group.
3	"(2) Appointment.—The members of the Work-
4	ing Group shall be appointed by the Secretary, except
5	that of the non-Federal public members under para-
6	graph (1)(B)—
7	"(A) one shall be appointed by the Speaker
8	of the House of Representatives; and
9	"(B) one shall be appointed by the majority
10	leader of the Senate.
11	"(3) Diversity of scientific perspectives.—
12	In making appointments under paragraph (2), the
13	Secretary, the Speaker of the House of Representa-
14	tives, and the majority leader of the Senate shall en-
15	sure that the non-Federal public members of the
16	Working Group represent a diversity of scientific per-
17	spectives.
18	"(4) Terms.—The non-Federal public members
19	of the Working Group shall each be appointed to serve
20	a 4-year term and may be reappointed at the end of
21	such term.
22	"(d) Meetings.—The Working Group shall meet as
23	often as necessary, as determined by the Secretary, but not
24	less than twice each year.

1	"(e) Applicability of FACA.—The Working Group
2	shall be treated as an advisory committee subject to the Fed-
3	eral Advisory Committee Act.
4	"(f) Reporting.—Not later than 24 months after the
5	date of enactment of this part, and every 24 months there-
6	after, the Working Group—
7	"(1) shall submit a report on its activities, in-
8	cluding an up-to-date summary under subsection
9	(b)(1) and any recommendations under subsection
10	(b)(4), to the Secretary, the Committee on Energy and
11	Commerce of the House of Representatives, and the
12	Committee on Health, Education, Labor and Pen-
13	sions of the Senate;
14	"(2) shall make each such report publicly avail-
15	able on the website of the Department of Health and
16	Human Services; and
17	"(3) shall allow any member of the Working
18	Group to include in any such report minority views.
19	"SEC. 39900-2. STRATEGIC PLAN.
20	"Not later than 3 years after the date of enactment
21	of this section, and every 5 years thereafter, the Secretary
22	shall submit to the Congress a strategic plan, informed by
23	the most recent summary under section 39900-1(b)(1), for
24	the conduct and support of Lyme disease and tick-borne dis-
25	ease research, including—

1	"(1) proposed budgetary requirements;
2	"(2) a plan for improving outcomes of Lyme dis-
3	ease and other tick-borne diseases, including progress
4	related to chronic or persistent symptoms and chronic
5	or persistent infection and co-infections;
6	"(3) a plan for improving diagnosis, treatment,
7	and prevention;
8	"(4) appropriate benchmarks to measure
9	progress on achieving the improvements described in
10	paragraphs (2) and (3); and
11	"(5) a plan to disseminate each summary under
12	section 39900–1(b)(1) and other relevant information
13	developed by the Working Group to the public, includ-
14	ing health care providers, public health departments,
15	and other relevant medical groups.".
16	(b) No Additional Authorization of Appropria-
17	Tions.—No additional funds are authorized to be appro-
18	priated for the purpose of carrying out this section and the
19	amendment made by this section, and this section and such
20	amendment shall be carried out using amounts otherwise
21	available for such purpose.

Union Calendar No. 142

114TH CONGRESS H. R. 6

[Report No. 114-190, Part I]

A BILL

To accelerate the discovery, development, and delivery of 21st century cures, and for other purposes.

July 7, 2015

Reported from the Committee on Energy and Commerce with an amendment

July 7, 2015

The Committee on Ways and Means discharged; committed to the Committee of the Whole House on the State of the Union and ordered to be printed