To establish a fair price negotiation program, protect the Medicare program from excessive price increases, and establish an out-of-pocket maximum for Medicare part D enrollees, and for other purposes.

IN THE HOUSE OF REPRESENTATIVES

September 19, 2019

Mr. Pallone (for himself, Mr. Neal, and Mr. Scott of Virginia) introduced the following bill; which was referred to the Committee on Energy and Commerce, and in addition to the Committees on Ways and Means, and Education and Labor, 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.

November --, 2019

Reported from the Committee on Ways and Means with an amendment

[Strike out all after the enacting clause and insert the part printed in italic]

[For text of introduced bill, see copy of bill as introduced on September 19, 2019]
A BILL

To establish a fair price negotiation program, protect the Medicare program from excessive price increases, and establish an out-of-pocket maximum for Medicare part D enrollees, and for other purposes.
Be it enacted by the Senate and House of Representatives of the United States of America in Congress assembled,

SECTION 1. SHORT TITLE; TABLE OF CONTENTS.

(a) SHORT TITLE.—This Act may be cited as the “Elijah E. Cummings Lower Drug Costs Now Act”.

(b) TABLE OF CONTENTS.—The table of contents is as follows:

Sec. 1. Short title; table of contents.

TITLE I—LOWERING PRICES THROUGH FAIR DRUG PRICE NEGOTIATION

Sec. 101. Providing for lower prices for certain high-priced single source drugs.
Sec. 102. Selected drug manufacturer excise tax imposed during noncompliance periods.

TITLE II—MEDICARE PARTS B AND D PRESCRIPTION DRUG INFLATION REBATES

Sec. 201. Medicare part B rebate by manufacturers.

TITLE III—PART D IMPROVEMENTS AND MAXIMUM OUT-OF-POCKET CAP FOR MEDICARE BENEFICIARIES

Sec. 301. Medicare part D benefit redesign.
Sec. 302. Allowing certain enrollees of prescription drugs plans and MA–PD plans under Medicare program to spread out cost-sharing under certain circumstances.
Sec. 303. Establishment of pharmacy quality measures under Medicare part D.

TITLE IV—PRESCRIPTION DRUG POLICIES FOR LOW-INCOME INDIVIDUALS

Sec. 401. Adjustments to Medicare part D cost-sharing reductions for low-income individuals.
Sec. 402. Dissemination to Medicare part D subsidy eligible individuals of information comparing premiums of certain prescription drug plans.
Sec. 403. Providing for intelligent assignment of certain subsidy eligible individuals auto-enrolled under Medicare prescription drug plans and MA–PD plans.
Sec. 404. Expanding eligibility for low-income subsidies under part D of the Medicare program.
Sec. 405. Automatic eligibility of certain low-income territorial residents for premium and cost-sharing subsidies under the Medicare program; Sunset of enhanced allotment program.
Sec. 406. Automatic qualification of certain Medicaid beneficiaries for premium and cost-sharing subsidies under part D of the Medicare program.
Sec. 407. Eliminating the resource requirement with respect to subsidy eligible individuals under part D of the Medicare program.

Sec. 408. Providing for certain rules regarding the treatment of eligible retirement plans in determining the eligibility of individuals for premium and cost-sharing subsidies under part D of the Medicare program.

TITLE V—DRUG PRICE TRANSPARENCY

Sec. 501. Drug price transparency.

TITLE I—LOWERING PRICES THROUGH FAIR DRUG PRICE NEGOTIATION

SEC. 101. PROVIDING FOR LOWER PRICES FOR CERTAIN HIGH-PRICED SINGLE SOURCE DRUGS.

(a) Program To Lower Prices for Certain High-Priced Single Source Drugs.—Title XI of the Social Security Act (42 U.S.C. 1301 et seq.) is amended by adding at the end the following new part:

“PART E—FAIR PRICE NEGOTIATION PROGRAM TO LOWER PRICES FOR CERTAIN HIGH-PRICED SINGLE SOURCE DRUGS

“SEC. 1191. ESTABLISHMENT OF PROGRAM.

“(a) In General.—The Secretary shall establish a Fair Price Negotiation Program (in this part referred to as the ‘program’). Under the program, with respect to each price applicability period, the Secretary shall—

“(1) publish a list of selected drugs in accordance with section 1192;
“(2) enter into agreements with manufacturers of
selected drugs with respect to such period, in accord-
ance with section 1193;

“(3) negotiate and, if applicable, renegotiate
maximum fair prices for such selected drugs, in ac-
cordance with section 1194; and

“(4) carry out the administrative duties de-
scribed in section 1196.

“(b) DEFINITIONS RELATING TO TIMING.—For pur-
poses of this part:

“(1) INITIAL PRICE APPLICABILITY YEAR.—The
term ‘initial price applicability year’ means a plan
year (beginning with plan year 2023) or, if agreed to
in an agreement under section 1193 by the Secretary
and manufacturer involved, a period of more than
one plan year (beginning on or after January 1,
2023).

“(2) PRICE APPLICABILITY PERIOD.—The term
‘price applicability period’ means, with respect to a
drug, the period beginning with the initial price ap-
plicability year with respect to which such drug is a
selected drug and ending with the last plan year dur-
ing which the drug is a selected drug.

“(3) SELECTED DRUG PUBLICATION DATE.—The
term ‘selected drug publication date’ means, with re-
spect to each initial price applicability year, April 15 of the plan year that begins 2 years prior to such year.

“(4) VOLUNTARY NEGOTIATION PERIOD.—The term ‘voluntary negotiation period’ means, with respect to an initial price applicability year with respect to a selected drug, the period—

“(A) beginning on the sooner of—

“(i) the date on which the manufacturer of the drug and the Secretary enter into an agreement under section 1193 with respect to such drug; or

“(ii) June 15 following the selected drug publication date with respect to such selected drug; and

“(B) ending on March 31 of the year that begins one year prior to the initial price applicability year.

“(c) OTHER DEFINITIONS.—For purposes of this part:

“(1) FAIR PRICE ELIGIBLE INDIVIDUAL.—The term ‘fair price eligible individual’ means, with respect to a selected drug—

“(A) in the case such drug is furnished or dispensed to the individual at a pharmacy or by a mail order service—
“(i) an individual who is enrolled under a prescription drug plan under part D of title XVIII or an MA–PD plan under part C of such title under which coverage is provided for such drug; and

“(ii) an individual who is enrolled under a group health plan or health insurance coverage offered in the group or individual market (as such terms are defined in section 2791 of the Public Health Service Act) with respect to which there is in effect an agreement with the Secretary under section 1197 with respect to such selected drug as so furnished or dispensed; and

“(B) in the case such drug is furnished or administered to the individual by a hospital, physician, or other provider of services or supplier—

“(i) an individual who is entitled to benefits under part A of title XVIII or enrolled under part B of such title if such selected drug is covered under the respective part; and

“(ii) an individual who is enrolled under a group health plan or health insur-
ance coverage offered in the group or individual market (as such terms are defined in section 2791 of the Public Health Service Act) with respect to which there is in effect an agreement with the Secretary under section 1197 with respect to such selected drug as so furnished or administered.

“(2) MAXIMUM FAIR PRICE.—The term ‘maximum fair price’ means, with respect to a plan year during a price applicability period and with respect to a selected drug (as defined in section 1192(c)) with respect to such period, the price published pursuant to section 1195 in the Federal Register for such drug and year.

“(3) AVERAGE INTERNATIONAL MARKET PRICE DEFINED.—

“(A) IN GENERAL.—The terms ‘average international market price’ and ‘AIM price’ mean, with respect to a drug, the average price (which shall be the net average price, if practicable, and volume-weighted, if practicable) for a unit (as defined in paragraph (4)) of the drug for sales of such drug (calculated across different dosage forms and strengths of the drug and not based on the specific formulation or package size
or package type), as computed (as of the date of
publication of such drug as a selected drug under
section 1192(a)) in all countries described in
clause (ii) of subparagraph (B) that are applica-
table countries (as described in clause (i) of such
subparagraph) with respect to such drug.

“(B) APPLICABLE COUNTRIES.—

“(i) IN GENERAL.—For purposes of
subparagraph (A), a country described in
clause (ii) is an applicable country de-
scribed in this clause with respect to a drug
if there is available an average price for
any unit for the drug for sales of such drug
in such country.

“(ii) COUNTRIES DESCRIBED.—For
purposes of this paragraph, the following
are countries described in this clause:

“(I) Australia.
“(II) Canada.
“(III) France.
“(IV) Germany.
“(V) Japan.
“(VI) The United Kingdom.

“(4) UNIT.—The term ‘unit’ means, with respect
to a drug, the lowest identifiable quantity (such as a
capsule or tablet, milligram of molecules, or grams) of the drug that is dispensed.

"SEC. 1192. SELECTION OF NEGOTIATION-ELIGIBLE DRUGS AS SELECTED DRUGS."

"(a) IN GENERAL.—Not later than the selected drug publication date with respect to an initial price applicability year, the Secretary shall select and publish in the Federal Register a list of—

"(1)(A) with respect to an initial price applicability year during the period beginning with 2023 and ending with 2027, at least 25 negotiation-eligible drugs described in subparagraphs (A) and (B), but not subparagraph (C), of subsection (d)(1) (or, with respect to an initial price applicability year during such period beginning after 2023, the maximum number (if such number is less than 25) of such negotiation-eligible drugs for the year) with respect to such year;

"(B) with respect to an initial price applicability year during the period beginning with 2028 and ending with 2032, at least 30 negotiation-eligible drugs described in subparagraphs (A) and (B), but not subparagraph (C), of subsection (d)(1) (or, with respect to an initial price applicability year during such period, the maximum number (if such number is
less than 30) of such negotiation-eligible drugs for the year) with respect to such year; and

“(C) with respect to an initial price applicability year beginning after 2032, at least 35 negotiation-eligible drugs described in subparagraphs (A) and (B), but not subparagraph (C), of subsection (d)(1) (or, with respect to an initial price applicability year during such period, the maximum number (if such number is less than 35) of such negotiation-eligible drugs for the year) with respect to such year;

“(2) all negotiation-eligible drugs described in subparagraph (C) of such subsection with respect to such year; and

“(3) all new-entrant negotiation-eligible drugs (as defined in subsection (g)(1)) with respect to such year.

Each drug published on the list pursuant to the previous sentence shall be subject to the negotiation process under section 1194 for the voluntary negotiation period with respect to such initial price applicability year (and the renegotiation process under such section as applicable for any subsequent year during the applicable price applicability period). In applying this subsection, any negotiation-eligible drug that is selected under this subsection for an initial price applicability year shall not count toward the required
minimum amount of drugs to be selected under paragraph (1) for any subsequent year, including such a drug so selected that is subject to renegotiation under section 1194.

“(b) SELECTION OF DRUGS.—In carrying out subsection (a)(1) the Secretary shall select for inclusion on the published list described in subsection (a) with respect to a price applicability period, the negotiation-eligible drugs that the Secretary projects will result in the greatest savings to the Federal Government or fair price eligible individuals during the price applicability period. In making this projection of savings for drugs for which there is an AIM price for a price applicability period, the savings shall be projected across different dosage forms and strengths of the drugs and not based on the specific formulation or package size or package type of the drugs, taking into consideration both the volume of drugs for which payment is made, to the extent such data is available, and the amount by which the net price for the drugs exceeds the AIM price for the drugs.

“(c) SELECTED DRUG.—For purposes of this part, each drug included on the list published under subsection (a) with respect to an initial price applicability year shall be referred to as a ‘selected drug’ with respect to such year and each subsequent plan year beginning before the first
plan year beginning after the date on which the Secretary
determines two or more drug products—

“(1) are approved or licensed (as applicable)—

“(A) under section 505(j) of the Federal
Food, Drug, and Cosmetic Act using such drug
as the listed drug; or

“(B) under section 351(k) of the Public
Health Service Act using such drug as the refer-
ence product; and

“(2) continue to be marketed.

“(d) NEGOTIATION-ELIGIBLE DRUG.—

“(1) IN GENERAL.—For purposes of this part,
the term ‘negotiation-eligible drug’ means, with re-
spect to the selected drug publication date with re-
pect to an initial price applicability year, a qual-
ifying single source drug, as defined in subsection (e),
that meets any of the following criteria:

“(A) COVERED PART D DRUGS.—The drug
is among the 125 covered part D drugs (as de-
defined in section 1860D–2(e)) for which there was
an estimated greatest net spending under parts
C and D of title XVIII, as determined by the
Secretary, during the most recent plan year
prior to such drug publication date for which
data are available.
“(B) OTHER DRUGS.—The drug is among the 125 drugs for which there was an estimated greatest net spending in the United States (including the 50 States, the District of Columbia, and the territories of the United States), as determined by the Secretary, during the most recent plan year prior to such drug publication date for which data are available.

“(C) INSULIN.—The drug is a qualifying single source drug described in subsection (e)(3).

“(2) CLARIFICATION.—In determining whether a qualifying single source drug satisfies any of the criteria described in paragraph (1), the Secretary shall, to the extent practicable, use data that is aggregated across dosage forms and strengths of the drug and not based on the specific formulation or package size or package type of the drug.

“(3) PUBLICATION.—Not later than the selected drug publication date with respect to an initial price applicability year, the Secretary shall publish in the Federal Register a list of negotiation-eligible drugs with respect to such selected drug publication date.

“(e) QUALIFYING SINGLE SOURCE DRUG.—For purposes of this part, the term ‘qualifying single source drug’ means any of the following:
“(1) DRUG PRODUCTS.—A drug that—

“(A) is approved under section 505(c) of the Federal Food, Drug, and Cosmetic Act and continues to be marketed pursuant to such approval; and

“(B) is not the listed drug for any drug that is approved and continues to be marketed under section 505(j) of such Act.

“(2) BIOLOGICAL PRODUCTS.—A biological product that—

“(A) is licensed under section 351(a) of the Public Health Service Act, including any product that has been deemed to be licensed under section 351 of such Act pursuant to section 7002(e)(4) of the Biologics Price Competition and Innovation Act of 2009, and continues to be marketed under section 351 of such Act; and

“(B) is not the reference product for any biological product that is licensed and continues to be marketed under section 351(k) of such Act.

“(3) INSULIN PRODUCT.—Notwithstanding paragraphs (1) and (2), any insulin product that is approved under subsection (c) or (j) of section 505 of the Federal Food, Drug, and Cosmetic Act or licensed under subsection (a) or (k) of section 351 of the Pub-
lic Health Service Act and continues to be marketed under such section 505 or 351, including any insulin product that has been deemed to be licensed under section 351(a) of the Public Health Service Act pursuant to section 7002(c)(4) of the Biologics Price Competition and Innovation Act of 2009 and continues to be marketed pursuant to such licensure.

For purposes of applying paragraphs (1) and (2), a drug or biological product that is marketed by the same sponsor or manufacturer (or an affiliate thereof or a cross-licensed producer or distributor) as the listed drug or reference product described in such respective paragraph shall not be taken into consideration.

“(f) INFORMATION ON INTERNATIONAL DRUG PRICES.—For purposes of determining which negotiation-eligible drugs to select under subsection (a) and, in the case of such drugs that are selected drugs, to determine the maximum fair price for such a drug and whether such maximum fair price should be renegotiated under section 1194, the Secretary shall use data relating to the AIM price with respect to such drug as available or provided to the Secretary and shall on an ongoing basis request from manufacturers of selected drugs information on the AIM price of such a drug.

“(g) NEW-ENTRANT NEGOTIATION-ELIGIBLE DRUGS.—
“(1) IN GENERAL.—For purposes of this part, the term ‘new-entrant negotiation-eligible drug’ means, with respect to the selected drug publication date with respect to an initial price applicability year, a qualifying single source drug—

“(A) that is first approved or licensed, as described in paragraph (1), (2), or (3) of subsection (e), as applicable, during the year preceding such selected drug publication date; and

“(B) that the Secretary determines under paragraph (2) is likely to be included as a negotiation-eligible drug with respect to the subsequent selected drug publication date.

“(2) DETERMINATION.—In the case of a qualifying single source drug that meets the criteria described in subparagraph (A) of paragraph (1), with respect to an initial price applicability year, if the wholesale acquisition cost at which such drug is first marketed in the United States is equal to or greater than the median household income (as determined according to the most recent data collected by the United States Census Bureau), the Secretary shall determine before the selected drug publication date with respect to the initial price applicability year, if the drug is likely to be included as a negotiation-eligible
drug with respect to the subsequent selected drug publication date, based on the projected spending under title XVIII or in the United States on such drug. For purposes of this paragraph the term ‘United States’ includes the 50 States, the District of Columbia, and the territories of the United States.

“SEC. 1193. MANUFACTURER AGREEMENTS.

“(a) In General.—For purposes of section 1191(a)(2), the Secretary shall enter into agreements with manufacturers of selected drugs with respect to a price applicability period, by not later than June 15 following the selected drug publication date with respect to such selected drug, under which—

“(1) during the voluntary negotiation period for the initial price applicability year for the selected drug, the Secretary and manufacturer, in accordance with section 1194, negotiate to determine (and, by not later than the last date of such period and in accordance with subsection (c), agree to) a maximum fair price for such selected drug of the manufacturer in order to provide access to such price—

“(A) to fair price eligible individuals who with respect to such drug are described in subparagraph (A) of section 1191(c)(1) and are furnished or dispensed such drug during, subject to
subparagraph (2), the price applicability period;

and

“(B) to hospitals, physicians, and other providers of services and suppliers with respect to fair price eligible individuals who with respect to such drug are described in subparagraph (B) of such section and are furnished or administered such drug during, subject to subparagraph (2), the price applicability period;

“(2) the Secretary and the manufacturer shall, in accordance with a process and during a period specified by the Secretary pursuant to rulemaking, renegotiate (and, by not later than the last date of such period and in accordance with subsection (c), agree to) the maximum fair price for such drug if the Secretary determines that there is a material change in any of the factors described in section 1194(d) relating to the drug, including changes in the AIM price for such drug, in order to provide access to such maximum fair price (as so renegotiated)—

“(A) to fair price eligible individuals who with respect to such drug are described in subparagraph (A) of section 1191(c)(1) and are furnished or dispensed such drug during any year during the price applicability period (beginning
after such renegotiation) with respect to such selected drug; and

“(B) to hospitals, physicians, and other providers of services and suppliers with respect to fair price eligible individuals who with respect to such drug are described in subparagraph (B) of such section and are furnished or administered such drug during any year described in subparagraph (A);

“(3) the maximum fair price (including as renegotiated pursuant to paragraph (2)), with respect to such a selected drug, shall be provided to fair price eligible individuals, who with respect to such drug are described in subparagraph (A) of section 1191(c)(1), at the pharmacy or by a mail order service at the point-of-sale of such drug;

“(4) the manufacturer, subject to subsection (d), submits to the Secretary, in a form and manner specified by the Secretary—

“(A) for the voluntary negotiation period for the price applicability period (and, if applicable, before any period of renegotiation specified pursuant to paragraph (2)) with respect to such drug all information that the Secretary requires to carry out the negotiation (or renegotiation
process) under this part, including information described in section 1192(f) and section 1194(d)(1); and

“(B) on an ongoing basis, information on changes in prices for such drug that would affect the AIM price for such drug or otherwise provide a basis for renegotiation of the maximum fair price for such drug pursuant to paragraph (2);

“(5) the manufacturer agrees that in the case the selected drug of a manufacturer is a drug described in subsection (c), the manufacturer will, in accordance with such subsection, make any payment required under such subsection with respect to such drug; and

“(6) the manufacturer complies with requirements imposed by the Secretary for purposes of administering the program, including with respect to the duties described in section 1196.

“(b) AGREEMENT IN EFFECT UNTIL DRUG IS NO LONGER A SELECTED DRUG.—An agreement entered into under this section shall be effective, with respect to a drug, until such drug is no longer considered a selected drug under section 1192(c).

“(c) SPECIAL RULE FOR CERTAIN SELECTED DRUGS WITHOUT AIM PRICE.—
“(1) IN GENERAL.—In the case of a selected drug
for which there is no AIM price available with respect
to the initial price applicability year for such drug
and for which an AIM price becomes available begin-
ning with respect to a subsequent plan year during
the price applicability period for such drug, if the
Secretary determines that the amount described in
paragraph (2)(A) for a unit of such drug is greater
than the amount described in paragraph (2)(B) for a
unit of such drug, then by not later than one year
after the date of such determination, the manufac-
turer of such selected drug shall pay to the Treasury
an amount equal to the product of—

“(A) the difference between such amount de-
scribed in paragraph (2)(A) for a unit of such
drug and such amount described in paragraph
(2)(B) for a unit of such drug; and

“(B) the number of units of such drug sold
in the United States, including the 50 States, the
District of Columbia, and the territories of the
United States, during the period described in
paragraph (2)(B).

“(2) AMOUNTS DESCRIBED.—

“(A) WEIGHTED AVERAGE PRICE BEFORE
AIM PRICE AVAILABLE.—For purposes of para-
graph (1), the amount described in this subparagraph for a selected drug described in such paragraph, is the amount equal to the weighted average manufacturer price (as defined in section 1927(k)(1)) for such dosage strength and form for the drug during the period beginning with the first plan year for which the drug is included on the list of negotiation-eligible drugs published under section 1192(d) and ending with the last plan year during the price applicability period for such drug with respect to which there is no AIM price available for such drug.

“(B) AMOUNT MULTIPLIER AFTER AIM PRICE AVAILABLE.—For purposes of paragraph (1), the amount described in this subparagraph for a selected drug described in such paragraph, is the amount equal to 200 percent of the AIM price for such drug with respect to the first plan year during the price applicability period for such drug with respect to which there is an AIM price available for such drug.

“(d) CONFIDENTIALITY OF INFORMATION.—Information submitted to the Secretary under this part by a manufacturer of a selected drug that is proprietary information of such manufacturer (as determined by the Secretary) may
be used only by the Secretary or disclosed to and used by
the Comptroller General of the United States or the Medi-
care Payment Advisory Commission for purposes of car-
rying out this part.

“(e) REGULATIONS.—

“(1) IN GENERAL.—The Secretary shall, pursu-
ant to rulemaking, specify, in accordance with para-
graph (2), the information that must be submitted
under subsection (a)(4).

“(2) INFORMATION SPECIFIED.—Information de-
scribed in paragraph (1), with respect to a selected
drug, shall include information on sales of the drug
(by the manufacturer of the drug or by another entity
under license or other agreement with the manufac-
turer, with respect to the sales of such drug, regardless
of the name under which the drug is sold) in any for-
eign country that is part of the AIM price. The Sec-
retary shall verify, to the extent practicable, such
sales from appropriate officials of the government of
the foreign country involved.

“(f) COMPLIANCE WITH REQUIREMENTS FOR ADMIN-
ISTRATION OF PROGRAM.—Each manufacturer with an
agreement in effect under this section shall comply with re-
quirements imposed by the Secretary or a third party with
a contract under section 1196(c)(1), as applicable, for purposes of administering the program.

“SEC. 1194. NEGOTIATION AND RENEGOTIATION PROCESS.

“(a) In General.—For purposes of this part, under an agreement under section 1193 between the Secretary and a manufacturer of a selected drug, with respect to the period for which such agreement is in effect and in accordance with subsections (b) and (c), the Secretary and the manufacturer—

“(1) shall during the voluntary negotiation period with respect to the initial price applicability year for such drug, in accordance with this section, negotiate a maximum fair price for such drug for the purpose described in section 1193(a)(1); and

“(2) as applicable pursuant to section 1193(a)(2) and in accordance with the process specified pursuant to such section, renegotiate such maximum fair price for such drug for the purpose described in such section.

“(b) Negotiating Methodology and Objective.—

“(1) In General.—The Secretary shall develop and use a consistent methodology for negotiations under subsection (a) that, in accordance with paragraph (2) and subject to paragraph (3), achieves the
lowest maximum fair price for each selected drug
while appropriately rewarding innovation.

“(2) PRIORITIZING FACTORS.—In considering
the factors described in subsection (d) in negotiating
(and, as applicable, renegotiating) the maximum fair
price for a selected drug, the Secretary shall, to the
extent practicable, consider all of the available factors
listed but shall prioritize the following factors:

“(A) RESEARCH AND DEVELOPMENT
COSTS.—The factor described in paragraph
(1)(A) of subsection (d).

“(B) MARKET DATA.—The factor described
in paragraph (1)(B) of such subsection.

“(C) UNIT COSTS OF PRODUCTION AND DIS-
TRIBUTION.—The factor described in paragraph
(1)(C) of such subsection.

“(D) COMPARISON TO EXISTING THERA-
PEUTIC ALTERNATIVES.—The factor described in
paragraph (2)(A) of such subsection.

“(3) REQUIREMENT.—

“(A) IN GENERAL.—In negotiating the max-
imum fair price of a selected drug, with respect
to an initial price applicability year for the se-
lected drug, and, as applicable, in renegotiating
the maximum fair price for such drug, with re-
spect to a subsequent year during the price applicability period for such drug, in the case that
the manufacturer of the selected drug offers under the negotiation or renegotiation, as applicable, a price for such drug that is not more than the target price described in subparagraph (B) for such drug for the respective year, the Secretary shall agree under such negotiation or renegotiation, respectively, to such offered price as the maximum fair price.

“(B) TARGET PRICE.—

“(i) IN GENERAL.—Subject to clause (ii), the target price described in this subparagraph for a selected drug with respect to a year, is the average price (which shall be the net average price, if practicable, and volume-weighted, if practicable) for a unit of such drug for sales of such drug, as computed (across different dosage forms and strengths of the drug and not based on the specific formulation or package size or package type of the drug) in the applicable country described in section 1191(c)(3)(B) with respect to such drug that, with respect to such year, has the lowest average price
for such drug as compared to the average prices (as so computed) of such drug with respect to such year in the other applicable countries described in such section with respect to such drug.

“(ii) SELECTED DRUGS WITHOUT AIM PRICE.—In applying this paragraph in the case of negotiating the maximum fair price of a selected drug for which there is no AIM price available with respect to the initial price applicability year for such drug, or, as applicable, renegotiating the maximum fair price for such drug with respect to a subsequent year during the price applicability period for such drug before the first plan year for which there is an AIM price available for such drug, the target price described in this subparagraph for such drug and respective year is the amount that is 80 percent of the average manufacturer price (as defined in section 1927(k)(1)) for such drug and year.

“(4) ANNUAL REPORT.—After the completion of each voluntary negotiation period, the Secretary shall submit to Congress a report on the maximum fair
prices negotiated (or, as applicable, renegotiated) for such period. Such report shall include information on how such prices so negotiated (or renegotiated) meet the requirements of this part, including the requirements of this subsection.

“(c) LIMITATION.—

“(1) IN GENERAL.—Subject to paragraph (2), the maximum fair price negotiated (including as renegotiated) under this section for a selected drug, with respect to each plan year during a price applicability period for such drug, shall not exceed 120 percent of the AIM price applicable to such drug with respect to such year.

“(2) SELECTED DRUGS WITHOUT AIM PRICE.—In the case of a selected drug for which there is no AIM price available with respect to the initial price applicability year for such drug, for each plan year during the price applicability period before the first plan year for which there is an AIM price available for such drug, the maximum fair price negotiated (including as renegotiated) under this section for the selected drug shall not exceed the amount equal to 85 percent of the average manufacturer price for the drug with respect to such year.
“(d) CONSIDERATIONS.—For purposes of negotiating and, as applicable, renegotiating (including for purposes of determining whether to renegotiate) the maximum fair price of a selected drug under this part with the manufacturer of the drug, the Secretary shall, consistent with subsection (b)(2), take into consideration the following factors:

“(1) MANUFACTURER-SPECIFIC INFORMATION.—

The following information, including as submitted by the manufacturer:

“(A) Research and development costs of the manufacturer for the drug and the extent to which the manufacturer has recouped research and development costs.

“(B) Market data for the drug, including the distribution of sales across different programs and purchasers and projected future revenues for the drug.

“(C) Unit costs of production and distribution of the drug.

“(D) Prior Federal financial support for novel therapeutic discovery and development with respect to the drug.

“(E) Data on patents and on existing and pending exclusivity for the drug.

“(F) National sales data for the drug.
“(G) Information on clinical trials for the
drug in the United States or in applicable coun-
tries described in section 1191(c)(3)(B).

“(2) INFORMATION ON ALTERNATIVE PROD-
UCTS.—The following information:

“(A) The extent to which the drug rep-
resents a therapeutic advance as compared to ex-
isting therapeutic alternatives and, to the extent
such information is available, the costs of such
existing therapeutic alternatives.

“(B) Information on approval by the Food
and Drug Administration of alternative drug
products.

“(C) Information on comparative effective-
ness analysis for such products, taking into con-
sideration the effects of such products on specific
populations, such as individuals with disabil-
ities, the elderly, terminally ill, children, and
other patient populations.

In considering information described in subpara-
graph (C), the Secretary shall not use evidence or
findings from comparative clinical effectiveness re-
search in a manner that treats extending the life of
an elderly, disabled, or terminally ill individual as of
lower value than extending the life of an individual
who is younger, nondisabled, or not terminally ill.

Nothing in the previous sentence shall affect the application or consideration of an AIM price for a selected drug.

“(3) FOREIGN SALES INFORMATION.—To the extent available on a timely basis, including as provided by a manufacturer of the selected drug or otherwise, information on sales of the selected drug in each of the countries described in section 1191(c)(3)(B).

“(4) ADDITIONAL INFORMATION.—Information submitted to the Secretary, in accordance with a process specified by the Secretary, by other parties that are affected by the establishment of a maximum fair price for the selected drug.

“(e) REQUEST FOR INFORMATION.—For purposes of negotiating and, as applicable, renegotiating (including for purposes of determining whether to renegotiate) the maximum fair price of a selected drug under this part with the manufacturer of the drug, with respect to a price applicability period, and other relevant data for purposes of this section—

“(1) the Secretary shall, not later than the selected drug publication date with respect to the initial price applicability year of such period, request drug pricing information from the manufacturer of such

...
selected drug, including information described in sub-
section (d)(1); and

“(2) by not later than October 1 following the se-
lected drug publication date, the manufacturer of such
selected drug shall submit to the Secretary such re-
quested information in such form and manner as the
Secretary may require.

The Secretary shall request, from the manufacturer or oth-
ers, such additional information as may be needed to carry
out the negotiation and renegotiation process under this sec-
tion.

“SEC. 1195. PUBLICATION OF MAXIMUM FAIR PRICES.

“(a) IN GENERAL.—With respect to an initial price
applicability year and selected drug with respect to such
year, not later than April 1 of the plan year prior to such
initial price applicability year, the Secretary shall publish
in the Federal Register the maximum fair price for such
drug negotiated under this part with the manufacturer of
such drug.

“(b) UPDATES.—

“(1) SUBSEQUENT YEAR MAXIMUM FAIR
PRICES.—For a selected drug, for each plan year sub-
sequent to the initial price applicability year for such
drug with respect to which an agreement for such
drug is in effect under section 1193, the Secretary shall publish in the Federal Register—

“(A) subject to subparagraph (B), the amount equal to the maximum fair price published for such drug for the previous year, increased by the annual percentage increase in the consumer price index for all urban consumers (all items; U.S. city average) as of September of such previous year; or

“(B) in the case the maximum fair price for such drug was renegotiated, for the first year for which such price as so renegotiated applies, such renegotiated maximum fair price.

“(2) PRICES NEGOTIATED AFTER DEADLINE.—In the case of a selected drug with respect to an initial price applicability year for which the maximum fair price is determined under this part after the date of publication under this section, the Secretary shall publish such maximum fair price in the Federal Register by not later than 30 days after the date such maximum price is so determined.

“SEC. 1196. ADMINISTRATIVE DUTIES; COORDINATION PROVISIONS.

“(a) ADMINISTRATIVE DUTIES.—
“(1) In general.—For purposes of section 1191, the administrative duties described in this section are the following:

“(A) The establishment of procedures (including through agreements with manufacturers under this part, contracts with prescription drug plans under part D of title XVIII and MA–PD plans under part C of such title, and agreements under section 1197 with group health plans and health insurance issuers of health insurance coverage offered in the individual or group market) under which the maximum fair price for a selected drug is provided to fair price eligible individuals, who with respect to such drug are described in subparagraph (A) of section 1191(c)(1), at pharmacies or by mail order service at the point-of-sale of the drug for the applicable price period for such drug and providing that such maximum fair price is used for determining cost-sharing under such plans or coverage for the selected drug.

“(B) The establishment of procedures (including through agreements with manufacturers under this part and contracts with hospitals, physicians, and other providers of services and
suppliers and agreements under section 1197 with group health plans and health insurance issuers of health insurance coverage offered in the individual or group market) under which, in the case of a selected drug furnished or administered by such a hospital, physician, or other provider of services or supplier to fair price eligible individuals (who with respect to such drug are described in subparagraph (B) of section 1191(c)(1)), the maximum fair price for the selected drug is provided to such hospitals, physicians, and other providers of services and suppliers (as applicable) with respect to such individuals and providing that such maximum fair price is used for determining cost-sharing under the respective part, plan, or coverage for the selected drug.

“(C) The establishment of procedures (including through agreements and contracts described in subparagraphs (A) and (B)) to ensure that, not later than 90 days after the dispensing of a selected drug to a fair price eligible individual by a pharmacy or mail order service, the pharmacy or mail order service is reimbursed for an amount equal to the difference between—
“(i) the lesser of—

“(I) the wholesale acquisition cost

of the drug;

“(II) the national average drug

acquisition cost of the drug; and

“(III) any other similar deter-

mination of pharmacy acquisition

costs of the drug, as determined by the

Secretary; and

“(ii) the maximum fair price for the
drug.

“(D) The establishment of procedures to en-

sure that the maximum fair price for a selected
drug is applied before—

“(i) any coverage or financial assist-

ance under other health benefit plans or

programs that provide coverage or financial

assistance for the purchase or provision of

prescription drug coverage on behalf of fair

price eligible individuals as the Secretary

may specify; and

“(ii) any other discounts.

“(E) The establishment of procedures to

enter into appropriate agreements and protocols

for the ongoing computation of AIM prices for
selected drugs, including, to the extent possible, to compute the AIM price for selected drugs and including by providing that the manufacturer of such a selected drug should provide information for such computation not later than 3 months after the first date of the voluntary negotiation period for such selected drug.

“(F) The establishment of procedures to compute and apply the maximum fair price across different strengths and dosage forms of a selected drug and not based on the specific formulation or package size or package type of the drug.

“(G) The establishment of procedures to negotiate and apply the maximum fair price in a manner that does not include any dispensing or similar fee.

“(H) The establishment of procedures to carry out the provisions of this part, as applicable, with respect to—

“(i) fair price eligible individuals who are enrolled under a prescription drug plan under part D of title XVIII or an MA–PD plan under part C of such title;
“(ii) fair price eligible individuals who are enrolled under a group health plan or health insurance coverage offered by a health insurance issuer in the individual or group market with respect to which there is an agreement in effect under section 1197; and

“(iii) fair price eligible individuals who are entitled to benefits under part A of title XVIII or enrolled under part B of such title.

“(I) The establishment of a negotiation process and renegotiation process in accordance with section 1194, including a process for acquiring information described in subsection (d) of such section and determining amounts described in subsection (b) of such section.

“(J) The provision of a reasonable dispute resolution mechanism to resolve disagreements between manufacturers, fair price eligible individuals, and the third party with a contract under subsection (c)(1).

“(2) MONITORING COMPLIANCE.—

“(A) IN GENERAL.—The Secretary shall monitor compliance by a manufacturer with the
terms of an agreement under section 1193, including by establishing a mechanism through which violations of such terms may be reported.

“(B) NOTIFICATION.—If a third party with a contract under subsection (c)(1) determines that the manufacturer is not in compliance with such agreement, the third party shall notify the Secretary of such noncompliance for appropriate enforcement under section 4192 of the Internal Revenue Code of 1986 or section 1198, as applicable.

“(b) COLLECTION OF DATA.—

“(1) FROM PRESCRIPTION DRUG PLANS AND MA–PD PLANS.—The Secretary may collect appropriate data from prescription drug plans under part D of title XVIII and MA–PD plans under part C of such title in a timeframe that allows for maximum fair prices to be provided under this part for selected drugs.

“(2) FROM HEALTH PLANS.—The Secretary may collect appropriate data from group health plans or health insurance issuers offering group or individual health insurance coverage in a timeframe that allows for maximum fair prices to be provided under this part for selected drugs.
“(c) CONTRACT WITH THIRD PARTIES.—

“(1) IN GENERAL.—The Secretary may enter into a contract with 1 or more third parties to administer the requirements established by the Secretary in order to carry out this part. At a minimum, the contract with a third party under the preceding sentence shall require that the third party—

“(A) receive and transmit information between the Secretary, manufacturers, and other individuals or entities the Secretary determines appropriate;

“(B) receive, distribute, or facilitate the distribution of funds of manufacturers to appropriate individuals or entities in order to meet the obligations of manufacturers under agreements under this part;

“(C) provide adequate and timely information to manufacturers, consistent with the agreement with the manufacturer under this part, as necessary for the manufacturer to fulfill its obligations under this part; and

“(D) permit manufacturers to conduct periodic audits, directly or through contracts, of the data and information used by the third party to
determine discounts for applicable drugs of the manufacturer under the program.

“(2) PERFORMANCE REQUIREMENTS.—The Secretary shall establish performance requirements for a third party with a contract under paragraph (1) and safeguards to protect the independence and integrity of the activities carried out by the third party under the program under this part.

“SEC. 1197. VOLUNTARY PARTICIPATION BY OTHER HEALTH PLANS.

“(a) AGREEMENT TO PARTICIPATE UNDER PROGRAM.—

“(1) IN GENERAL.—Subject to paragraph (2), under the program under this part the Secretary shall be treated as having in effect an agreement with a group health plan or health insurance issuer offering health insurance coverage (as such terms are defined in section 2791 of the Public Health Service Act), with respect to a price applicability period and a selected drug with respect to such period—

“(A) with respect to such selected drug furnished or dispensed at a pharmacy or by mail order service if coverage is provided under such plan or coverage during such period for such selected drug as so furnished or dispensed; and
“(B) with respect to such selected drug furnished or administered by a hospital, physician, or other provider of services or supplier if coverage is provided under such plan or coverage during such period for such selected drug as so furnished or administered.

“(2) OPTING OUT OF AGREEMENT.—The Secretary shall not be treated as having in effect an agreement under the program under this part with a group health plan or health insurance issuer offering health insurance coverage with respect to a price applicability period and a selected drug with respect to such period if such a plan or issuer affirmatively elects, through a process specified by the Secretary, not to participate under the program with respect to such period and drug.

“(b) PUBLICATION OF ELECTION.—With respect to each price applicability period and each selected drug with respect to such period, the Secretary and the Secretary of Labor and the Secretary of the Treasury, as applicable, shall make public a list of each group health plan and each issuer of health insurance coverage, with respect to which coverage is provided under such plan or coverage for such drug, that has elected under subsection (a) not to partici-
pate under the program with respect to such period and drug.

“SEC. 1198. CIVIL MONETARY PENALTY.

“(a) Violations Relating To Offering Of Maximum Fair Price.—Any manufacturer of a selected drug that has entered into an agreement under section 1193, with respect to a plan year during the price applicability period for such drug, that does not provide access to a price that is not more than the maximum fair price (or a lesser price) for such drug for such year—

“(1) to a fair price eligible individual who with respect to such drug is described in subparagraph (A) of section 1191(c)(1) and who is furnished or dispensed such drug during such year; or

“(2) to a hospital, physician, or other provider of services or supplier with respect to fair price eligible individuals who with respect to such drug is described in subparagraph (B) of such section and is furnished or administered such drug by such hospital, physician, or provider or supplier during such year; shall be subject to a civil monetary penalty equal to ten times the amount equal to the difference between the price for such drug made available for such year by such manufacturer with respect to such individual or hospital, physi-
cian, provider, or supplier and the maximum fair price for such drug for such year.

“(b) Violations of Certain Terms of Agreement.—Any manufacturer of a selected drug that has entered into an agreement under section 1193, with respect to a plan year during the price applicability period for such drug, that is in violation of a requirement imposed pursuant to section 1193(a)(6) shall be subject to a civil monetary penalty of not more than $1,000,000 for each such violation.

“(c) Application.—The provisions of section 1128A (other than subsections (a) and (b)) shall apply to a civil monetary penalty under this section in the same manner as such provisions apply to a penalty or proceeding under section 1128A(a).


“(a) Paperwork Reduction Act.—Chapter 35 of title 44, United States Code, shall not apply to data collected under this part.

“(b) National Academy of Medicine Study.—Not later than December 31, 2025, the National Academy of Medicine shall conduct a study, and submit to Congress a report, on recommendations for improvements to the program under this part, including the determination of the limits applied under section 1194(c).
“(c) **MEDPAC STUDY.**—Not later than December 31, 2025, the Medicare Payment Advisory Commission shall conduct a study, and submit to Congress a report, on the program under this part with respect to the Medicare program under title XVIII, including with respect to the effect of the program on individuals entitled to benefits or enrolled under such title.

“(d) **LIMITATION ON JUDICIAL REVIEW.**—The following shall not be subject to judicial review:

“(1) The selection of drugs for publication under section 1192(a).

“(2) The determination of whether a drug is a negotiation-eligible drug under section 1192(d).

“(3) The determination of the maximum fair price of a selected drug under section 1194.

“(4) The determination of units of a drug for purposes of section 1191(c)(3).

“(e) **COORDINATION.**—In carrying out this part with respect to group health plans or health insurance coverage offered in the group market that are subject to oversight by the Secretary of Labor or the Secretary of the Treasury, the Secretary of Health and Human Services shall coordinate with such respective Secretary.

“(f) **DATA SHARING.**—The Secretary shall share with the Secretary of the Treasury such information as is nec-
(b) Application of Maximum Fair Prices and Conforming Amendments.—

(1) Under Medicare.—

(A) Application to Payments under Part B.—Section 1847A(b)(1)(B) of the Social Security Act (42 U.S.C. 1395w–3a(b)(1)(B)) is amended by inserting “or in the case of such a drug or biological that is a selected drug (as defined in section 1192(c)), with respect to a price applicability period (as defined in section 1191(b)(2)), 106 percent of the maximum fair price (as defined in section 1191(c)(2) applicable for such drug and a plan year during such period” after “paragraph (4)”.

(B) Exception to Part D Non-Interference.—Section 1860D–11(i) of the Social Security Act (42 U.S.C. 1395w–111(i)) is amended by inserting “, except as provided under part E of title XI” after “the Secretary”.

(C) Application as Negotiated Price under Part D.—Section 1860D–2(d)(1) of the Social Security Act (42 U.S.C. 1395w–102(d)(1)) is amended—
(i) in subparagraph (B), by inserting
“, subject to subparagraph (D),” after “negotiated prices”; and
(ii) by adding at the end the following new subparagraph:

“(D) APPLICATION OF MAXIMUM FAIR PRICE
FOR SELECTED DRUGS.—In applying this section, in the case of a covered part D drug that
is a selected drug (as defined in section 1192(c)),
with respect to a price applicability period (as
defined in section 1191(b)(2)), the negotiated
prices used for payment (as described in this
subsection) shall be the maximum fair price (as
defined in section 1191(c)(2)) for such drug and
for each plan year during such period.”.

(D) INFORMATION FROM PRESCRIPTION
DRUG PLANS AND MA–PD PLANS REQUIRED.—

(i) PRESCRIPTION DRUG PLANS.—Section
1860D–12(b) of the Social Security Act
(42 U.S.C. 1395w–112(b)) is amended by
adding at the end the following new para-
graph:

“(8) PROVISION OF INFORMATION RELATED TO
MAXIMUM FAIR PRICES.—Each contract entered into
with a PDP sponsor under this part with respect to
a prescription drug plan offered by such sponsor shall require the sponsor to provide information to the Secretary as requested by the Secretary in accordance with section 1196(b).”.

(ii) MA–PD PLANS.—Section 1857(f)(3) of the Social Security Act (42 U.S.C. 1395w–27(f)(3)) is amended by adding at the end the following new subparagraph:

“(E) PROVISION OF INFORMATION RELATED TO MAXIMUM FAIR PRICES.—Section 1860D–12(b)(8).”.

(2) UNDER GROUP HEALTH PLANS AND HEALTH INSURANCE COVERAGE.—

(A) PHSA.—Part A of title XXVII of the Public Health Service Act is amended by inserting after section 2729 the following new section:

“SEC. 2729A. FAIR PRICE DRUG NEGOTIATION PROGRAM AND APPLICATION OF MAXIMUM FAIR PRICES.

“(a) IN GENERAL.—In the case of a group health plan or health insurance issuer offering health insurance coverage that is treated under section 1197 of the Social Security Act as having in effect an agreement with the Secretary under the Fair Price Drug Negotiation Program under part
E of title XI of such Act, with respect to a price applicability period (as defined in section 1191(b) of such Act) and a selected drug (as defined in section 1192(c) of such Act) with respect to such period with respect to which coverage is provided under such plan or coverage—

“(1) the provisions of such part shall apply to the plans or coverage offered by such plan or issuer, and to the individuals enrolled under such plans or coverage, during such period, with respect to such selected drug, in the same manner as such provisions apply to prescription drug plans and MA–PD plans, and to individuals enrolled under such prescription drug plans and MA–PD plans;

“(2) the plan or issuer shall apply any cost-sharing responsibilities under such plan or coverage, with respect to such selected drug, by substituting the maximum fair price negotiated under such part for such drug in lieu of the contracted rate under such plan or coverage for such selected drug; and

“(3) the Secretary shall apply the provisions of such part to such plan, issuer, and coverage, and such individuals so enrolled in such plans.

“(b) Notification Regarding Nonparticipation in Fair Drug Price Negotiation Program.—A group health plan or a health insurance issuer offering group or
individual health insurance coverage shall publicly disclose in a manner and in accordance with a process specified by the Secretary any election made under section 1197 of the Social Security Act by the plan or issuer to not participate in the Fair Drug Price Negotiation Program under part E of title XI of such Act with respect to a selected drug (as defined in section 1192(c) of such Act) for which coverage is provided under such plan or coverage before the beginning of the plan year for which such election was made.”.

(B) ERISA.—

(i) IN GENERAL.—Subpart B of part 7 of subtitle B of title I of the Employee Retirement Income Security Act of 1974 (29 U.S.C. 1181 et. seq.) is amended by adding at the end the following new section:

“SEC. 716. FAIR PRICE DRUG NEGOTIATION PROGRAM AND APPLICATION OF MAXIMUM FAIR PRICES.

“(a) IN GENERAL.—In the case of a group health plan or health insurance issuer offering group health insurance coverage that is treated under section 1197 of the Social Security Act as having in effect an agreement with the Secretary under the Fair Price Drug Negotiation Program under part E of title XI of such Act, with respect to a price applicability period (as defined in section 1191(b) of such
Act) and a selected drug (as defined in section 1192(c) of such Act) with respect to such period with respect to which coverage is provided under such plan or coverage—

“(1) the provisions of such part shall apply to the plans or coverage offered by such plan or issuer, and to the individuals enrolled under such plans or coverage, during such period, with respect to such selected drug, in the same manner as such provisions apply to prescription drug plans and MA–PD plans, and to individuals enrolled under such prescription drug plans and MA–PD plans;

“(2) the plan or issuer shall apply any cost-sharing responsibilities under such plan or coverage, with respect to such selected drug, by substituting the maximum fair price negotiated under such part for such drug in lieu of the contracted rate under such plan or coverage for such selected drug; and

“(3) the Secretary shall apply the provisions of such part to such plan, issuer, and coverage, and such individuals so enrolled in such plans.

“(b) Notification Regarding Nonparticipation in Fair Drug Price Negotiation Program.—A group health plan or a health insurance issuer offering group health insurance coverage shall publicly disclose in a manner and in accordance with a process specified by the Sec-
retary any election made under section 1197 of the Social
Security Act by the plan or issuer to not participate in
the Fair Drug Price Negotiation Program under part E
of title XI of such Act with respect to a selected drug (as
defined in section 1192(c) of such Act) for which coverage
is provided under such plan or coverage before the begin-
ing of the plan year for which such election was made.”.

(ii) CLERICAL AMENDMENT.—The table
of sections for part 7 of subtitle B of title
I of the Employee Retirement Income Secu-
rit y Act of 1974 is amended by adding at
the end the following:

“Sec. 716. Fair Price Drug Negotiation Program and application of maximum
fair prices.”.

(C) IRC.—

(i) IN GENERAL.—Subchapter B of
chapter 100 of the Internal Revenue Code of
1986 is amended by adding at the end the
following new section:

“SEC. 9816. FAIR PRICE DRUG NEGOTIATION PROGRAM AND
APPLICATION OF MAXIMUM FAIR PRICES.

“(a) IN GENERAL.—In the case of a group health plan
that is treated under section 1197 of the Social Security
Act as having in effect an agreement with the Secretary
under the Fair Price Drug Negotiation Program under part
E of title XI of such Act, with respect to a price applica-
bility period (as defined in section 1191(b) of such Act) and
a selected drug (as defined in section 1192(c) of such Act)
with respect to such period with respect to which coverage
is provided under such plan—
“(1) the provisions of such part shall apply, as
applicable—
“(A) if coverage of such selected drug is pro-
vided under such plan if the drug is furnished
or dispensed at a pharmacy or by a mail order
service, to the plan, and to the individuals en-
rolled under such plan during such period, with
respect to such selected drug, in the same manner
as such provisions apply to prescription drug
plans and MA–PD plans, and to individuals en-
rolled under such prescription drug plans and
MA–PD plans during such period; and
“(B) if coverage of such selected drug is pro-
vided under such plan if the drug is furnished
or administered by a hospital, physician, or
other provider of services or supplier, to the
plan, to the individuals enrolled under such
plan, and to hospitals, physicians, and other
providers of services and suppliers during such
period, with respect to such drug in the same
manner as such provisions apply to the Sec-
retary, to individuals entitled to benefits under part A of title XVIII or enrolled under part B of such title, and to hospitals, physicians, and other providers and suppliers participating under title XVIII during such period;

“(2) the plan shall apply any cost-sharing responsibilities under such plan, with respect to such selected drug, by substituting an amount not more than the maximum fair price negotiated under such part E of title XI for such drug in lieu of the drug price upon which the cost-sharing would have otherwise applied; and

“(3) the Secretary shall apply the provisions of such part E to such plan and such individuals so enrolled in such plan.

“(b) Notification Regarding Nonparticipation in Fair Drug Price Negotiation Program.—A group health plan shall publicly disclose in a manner and in accordance with a process specified by the Secretary any election made under section 1197 of the Social Security Act by the plan to not participate in the Fair Drug Price Negotiation Program under part E of title XI of such Act with respect to a selected drug (as defined in section 1192(c) of such Act) for which coverage is provided under such plan.
before the beginning of the plan year for which such election was made.”

(ii) APPLICATION TO RETIREE AND CERTAIN SMALL GROUP HEALTH PLANS.—

Section 9831(a)(2) of the Internal Revenue Code of 1986 is amended by inserting “other than with respect to section 9816,” before “any group health plan”.

(iii) CLERICAL AMENDMENT.—The table of sections for subchapter B of chapter 100 of such Code is amended by adding at the end the following new item:

“Sec. 9816. Fair Price Drug Negotiation Program and application of maximum fair prices.”.

SEC. 102. SELECTED DRUG MANUFACTURER EXCISE TAX IMPOSED DURING NONCOMPLIANCE PERIODS.

(a) IN GENERAL.—Subchapter E of chapter 32 of the Internal Revenue Code of 1986 is amended by adding at the end the following new section:

“SEC. 4192. SELECTED DRUGS DURING NONCOMPLIANCE PERIODS.

“(a) IN GENERAL.—There is hereby imposed on the sale by the manufacturer, producer, or importer of any selected drug during a day described in subsection (b) a tax in an amount such that the applicable percentage is equal to the ratio of—
“(1) such tax, divided by

“(2) the sum of such tax and the price for which

so sold.

“(b) NONCOMPLIANCE PERIODS.—A day is described

in this subsection with respect to a selected drug if it is

a day during one of the following periods:

“(1) The period beginning on the June 16th im-

mediately following the selected drug publication date

and ending on the first date during which the manu-

ufacturer of the drug has in place an agreement de-

scribed in subsection (a) of section 1193 of the Social

Security Act with respect to such drug.

“(2) The period beginning on the April 1st im-

mediately following the June 16th described in para-

graph (1) and ending on the first date during which

the manufacturer of the drug has agreed to a max-

imum fair price under such agreement.

“(3) In the case of a selected drug with respect

to which the Secretary of Health and Human Services

has specified a renegotiation period under such agree-

ment, the period beginning on the first date after the

last date of such renegotiation period and ending on

the first date during which the manufacturer of the

drug has agreed to a renegotiated maximum fair

price under such agreement.
“(4) With respect to information that is required to be submitted to the Secretary of Health and Human Services under such agreement, the period beginning on the date on which such Secretary certifies that such information is overdue and ending on the date that such information is so submitted.

“(5) In the case of a selected drug with respect to which a payment is due under subsection (c) of such section 1193, the period beginning on the date on which the Secretary of Health and Human Services certifies that such payment is overdue and ending on the date that such payment is made in full.

“(c) APPLICABLE PERCENTAGE.—For purposes of this section, the term ‘applicable percentage’ means—

“(1) in the case of sales of a selected drug during the first 90 days described in subsection (b) with respect to such drug, 65 percent,

“(2) in the case of sales of such drug during the 91st day through the 180th day described in subsection (b) with respect to such drug, 75 percent,

“(3) in the case of sales of such drug during the 181st day through the 270th day described in subsection (b) with respect to such drug, 85 percent, and

“(4) in the case of sales of such drug during any subsequent day, 95 percent.
“(d) SELECTED DRUG.—For purposes of this section—

“(1) IN GENERAL.—The term ‘selected drug’ means any selected drug (within the meaning of section 1192 of the Social Security Act) which is manufactured or produced in the United States or entered into the United States for consumption, use, or warehousing.

“(2) UNITED STATES.—The term ‘United States’ has the meaning given such term by section 4612(a)(4).

“(3) COORDINATION WITH RULES FOR POSSESSIONS OF THE UNITED STATES.—Rules similar to the rules of paragraphs (2) and (4) of section 4132(c) shall apply for purposes of this section.

“(e) OTHER DEFINITIONS.—For purposes of this section, the terms ‘selected drug publication date’ and ‘maximum fair price’ have the meaning given such terms in section 1191 of the Social Security Act.

“(f) ANTI-ABUSE RULE.—In the case of a sale which was timed for the purpose of avoiding the tax imposed by this section, the Secretary may treat such sale as occurring during a day described in subsection (b).”.

(b) NO DEDUCTION FOR EXCISE TAX PAYMENTS.—

Section 275 of the Internal Revenue Code of 1986 is amend-
ed by adding “or by section 4192” before the period at the
end of subsection (a)(6).

(c) CONFORMING AMENDMENTS.—

(1) Section 4221(a) of the Internal Revenue Code
of 1986 is amended by inserting “or 4192” after “sec-
tion 4191”.

(2) Section 6416(b)(2) of such Code is amended
by inserting “or 4192” after “section 4191”.

(d) CLERICAL AMENDMENTS.—

(1) The heading of subchapter E of chapter 32
of the Internal Revenue Code of 1986 is amended by
striking “Medical Devices” and inserting
“Other Medical Products”.

(2) The table of subchapters for chapter 32 of
such Code is amended by striking the item relating to
subchapter E and inserting the following new item:
“subchapter E. Other Medical Products”.

(3) The table of sections for subchapter E of
chapter 32 of such Code is amended by adding at the
end the following new item:
“Sec. 4192. Selected drugs during noncompliance periods.”.

(e) EFFECTIVE DATE.—The amendments made by this
section shall apply to sales after the date of the enactment
of this Act.
TITLE II—MEDICARE PARTS B AND D PRESCRIPTION DRUG INFLATION REBATES

SEC. 201. MEDICARE PART B REBATE BY MANUFACTURERS.

(a) In General.—Section 1834 of the Social Security Act (42 U.S.C. 1395m) is amended by adding at the end the following new subsection:

“(x) Rebate by Manufacturers for Single Source Drugs With Prices Increasing Faster Than Inflation.—

“(1) Requirements.—

“(A) Secretarial Provision of Information.—Not later than 6 months after the end of each calendar quarter beginning on or after July 1, 2021, the Secretary shall, for each part B rebatable drug, report to each manufacturer of such part B rebatable drug the following for such calendar quarter:

“(i) Information on the total number of units of the billing and payment code described in subparagraph (A)(i) of paragraph (3) with respect to such drug and calendar quarter.

“(ii) Information on the amount (if any) of the excess average sales price in-
crease described in subparagraph (A)(ii) of such paragraph for such drug and calendar quarter.

“(iii) The rebate amount specified under such paragraph for such part B rebatable drug and calendar quarter.

“(B) MANUFACTURER REQUIREMENT.—For each calendar quarter beginning on or after July 1, 2021, the manufacturer of a part B rebatable drug shall, for such drug, not later than 30 days after the date of receipt from the Secretary of the information described in subparagraph (A) for such calendar quarter, provide to the Secretary a rebate that is equal to the amount specified in paragraph (3) for such drug for such calendar quarter.

“(2) PART B REBATABLE DRUG DEFINED.—

“(A) IN GENERAL.—In this subsection, the term ‘part B rebatable drug’ means a single source drug or biological (as defined in subparagraph (D) of section 1847A(c)(6)), including a biosimilar biological product (as defined in subparagraph (H) of such section), paid for under this part, except such term shall not include such a drug or biological—
“(i) if the average total allowed charges for a year per individual that uses such a drug or biological, as determined by the Secretary, are less than, subject to subparagraph (B), $100; or

“(ii) that is a vaccine described in subparagraph (A) or (B) of section 1861(s)(10).

“(B) INCREASE.—The dollar amount applied under subparagraph (A)(i)—

“(i) for 2022, shall be the dollar amount specified under such subparagraph for 2021, increased by the percentage increase in the consumer price index for all urban consumers (United States city average) for the 12 month period ending with June of the previous year; and

“(ii) for a subsequent year, shall be the dollar amount specified in this clause (or clause (i)) for the previous year, increased by the percentage increase in the consumer price index for all urban consumers (United States city average) for the 12 month period ending with June of the previous year.
Any dollar amount specified under this subpara-
graph that is not a multiple of $10 shall be
rounded to the nearest multiple of $10.

“(3) REBATE AMOUNT.—

“(A) IN GENERAL.—For purposes of para-
graph (1), the amount specified in this para-
graph for a part B rebatable drug assigned to a
billing and payment code for a calendar quarter
is, subject to paragraph (4), the amount equal to
the product of—

“(i) subject to subparagraphs (B) and

(G), the total number of units of the billing
and payment code for such part B rebatable
drug furnished under this part during the
calendar quarter; and

“(ii) the amount (if any) by which—

“(I) the payment amount under

subparagraph (B) or (C) of section

1847A(b)(1), as applicable, for such
part B rebatable drug during the cal-
endar quarter; exceeds

“(II) the inflation-adjusted pay-
ment amount determined under sub-
paragraph (C) for such part B
rebatable drug during the calendar quarter.

“(B) EXCLUDED UNITS.—For purposes of subparagraph (A)(i), the total number of units of the billing and payment code for each part B rebatable drug furnished during a calendar quarter shall not include—

“(i) units packaged into the payment for a procedure or service under section 1833(t) or under section 1833(i) (instead of separately payable under such respective section);

“(ii) units included under the single payment system for renal dialysis services under section 1881(b)(14); or

“(iii) units of a part B rebatable drug of a manufacturer furnished to an individual, if such manufacturer, with respect to the furnishing of such units of such drug, provides for discounts under section 340B of the Public Health Service Act or for rebates under section 1927.

“(C) DETERMINATION OF INFLATION-ADJUSTED PAYMENT AMOUNT.—The inflation-adjusted payment amount determined under this
subparagraph for a part B rebatable drug for a
calendar quarter is—

“(i) the payment amount for the bill-
ing and payment code for such drug in the
payment amount benchmark quarter (as de-
defined in subparagraph (D)); increased by

“(ii) the percentage by which the rebate
period CPI–U (as defined in subparagraph
(F)) for the calendar quarter exceeds the
benchmark period CPI–U (as defined in
subparagraph (E)).

“(D) PAYMENT AMOUNT BENCHMARK QUAR-
ter.—The term ‘payment amount benchmark
quarter’ means the calendar quarter beginning
January 1, 2016.

“(E) BENCHMARK PERIOD CPI–U.—The
term ‘benchmark period CPI–U’ means the con-
sumer price index for all urban consumers
(United States city average) for July 2015.

“(F) REBATE PERIOD CPI–U.—The term
‘rebate period CPI–U’ means, with respect to a
calendar quarter described in subparagraph (C),
the greater of the benchmark period CPI–U and
the consumer price index for all urban con-
sumers (United States city average) for the first
month of the calendar quarter that is two calendar quarters prior to such described calendar quarter.

“(G) COUNTING UNITS.—

“(i) CUT-OFF PERIOD TO COUNT UNITS.—For purposes of subparagraph (A)(i), subject to clause (ii), to count the total number of billing units for a part B rebatable drug for a quarter, the Secretary may use a cut-off period in order to exclude from such total number of billing units for such quarter claims for services furnished during such quarter that were not processed at an appropriate time prior to the end of the cut-off period.

“(ii) COUNTING UNITS FOR CLAIMS PROCESSED AFTER CUT-OFF PERIOD.—If the Secretary uses a cut-off period pursuant to clause (i), in the case of units of a part B rebatable drug furnished during a quarter but pursuant to application of such cut-off period excluded for purposes of subparagraph (A)(i) from the total number of billing units for the drug for such quarter, the Secretary shall count such units of such
drug so furnished in the total number of billing units for such drug for a subsequent quarter, as the Secretary determines appropriate.

“(4) SPECIAL TREATMENT OF CERTAIN DRUGS AND EXEMPTION.—

“(A) SUBSEQUENTLY APPROVED DRUGS.— Subject to subparagraph (B), in the case of a part B rebatable drug first approved or licensed by the Food and Drug Administration after July 1, 2015, clause (i) of paragraph (3)(C) shall be applied as if the term ‘payment amount benchmark quarter’ were defined under paragraph (3)(D) as the third full calendar quarter after the day on which the drug was first marketed and clause (ii) of paragraph (3)(C) shall be applied as if the term ‘benchmark period CPI–U’ were defined under paragraph (3)(E) as if the reference to ‘July 2015’ under such paragraph were a reference to ‘the first month of the first full calendar quarter after the day on which the drug was first marketed’.

“(B) TIMELINE FOR PROVISION OF REBATES FOR SUBSEQUENTLY APPROVED DRUGS.—In the case of a part B rebatable drug first approved or
licensed by the Food and Drug Administration after July 1, 2015, paragraph (1)(B) shall be applied as if the reference to ‘July 1, 2021’ under such paragraph were a reference to the later of the 6th full calendar quarter after the day on which the drug was first marketed or July 1, 2021.

“(C) EXEMPTION FOR SHORTAGES.—The Secretary may reduce or waive the rebate amount under paragraph (1)(B) with respect to a part B rebateable drug that is described as currently in shortage on the shortage list in effect under section 506E of the Federal Food, Drug, and Cosmetic Act or in the case of other exigent circumstances, as determined by the Secretary.

“(D) SELECTED DRUGS.—In the case of a part B rebateable drug that is a selected drug (as defined in section 1192(c)) for a price applicability period (as defined in section 1191(b)(2)) and is determined (pursuant to such section 1192(c)) to no longer be a selected drug, for each applicable year beginning after the price applicability period with respect to such drug, clause (i) of paragraph (3)(C) shall be applied as if the term ‘payment amount benchmark quarter’ were
defined under paragraph (3)(D) as the calendar quarter beginning January 1 of the last year beginning during such price applicability period with respect to such selected drug and clause (ii) of paragraph (3)(C) shall be applied as if the term ‘benchmark period CPI–U’ were defined under paragraph (3)(E) as if the reference to ‘July 2015’ under such paragraph were a reference to the July of the year preceding such last year.

“(5) APPLICATION TO BENEFICIARY COINSURANCE.—In the case of a part B rebatable drug, if the payment amount for a quarter exceeds the inflation adjusted payment for such quarter—

“(A) in computing the amount of any coinsurance applicable under this title to an individual with respect to such drug, the computation of such coinsurance shall be based on the inflation-adjusted payment amount determined under paragraph (3)(C) for such part B rebatable drug; and

“(B) the amount of such coinsurance is equal to 20 percent of such inflation-adjusted payment amount so determined.
“(6) Rebate deposits.—Amounts paid as rebates under paragraph (1)(B) shall be deposited into the Federal Supplementary Medical Insurance Trust Fund established under section 1841.

“(7) Civil money penalty.—If a manufacturer of a part B rebatable drug has failed to comply with the requirements under paragraph (1)(B) for such drug for a calendar quarter, the manufacturer shall be subject to, in accordance with a process established by the Secretary pursuant to regulations, a civil money penalty in an amount equal to at least 125 percent of the amount specified in paragraph (3) for such drug for such calendar quarter. The provisions of section 1128A (other than subsections (a) (with respect to amounts of penalties or additional assessments) and (b)) shall apply to a civil money penalty under this paragraph in the same manner as such provisions apply to a penalty or proceeding under section 1128A(a).

“(8) Study and report.—

“(A) Study.—The Secretary shall conduct a study of the feasibility of and operational issues involved with the following:
“(i) Including multiple source drugs
(as defined in section 1847A(c)(6)(C)) in
the rebate system under this subsection.

“(ii) Including drugs and biologicals
paid for under MA plans under part C in
the rebate system under this subsection.

“(iii) Including drugs excluded under
paragraph (2)(A) and units of the billing
and payment code of the drugs excluded
under paragraph (3)(B) in the rebate sys-
tem under this subsection.

“(B) REPORT.—Not later than 3 years after
the date of the enactment of this subsection, the
Secretary shall submit to Congress a report on
the study conducted under subparagraph (A).

“(9) APPLICATION TO MULTIPLE SOURCE
DRUGS.—The Secretary may, based on the report sub-
mitted under paragraph (8) and pursuant to rule-
making, apply the provisions of this subsection to
multiple source drugs (as defined in section
1847A(c)(6)(C)), including, for purposes of deter-
mining the rebate amount under paragraph (3), by
calculating manufacturer-specific average sales prices
for the benchmark period and the rebate period.”.
(b) AMOUNTS PAYABLE; COST-SHARING.—Section 1833 of the Social Security Act (42 U.S.C. 1395l) is amended—

(1) in subsection (a)—

(A) in paragraph (1)—

(i) in subparagraph (S), by striking “with respect to” and inserting “subject to subparagraph (DD), with respect to”;

(ii) by striking “and (CC)” and inserting “(CC)”;

(iii) by inserting before the semicolon at the end the following: “, and (DD) with respect to a part B rebatable drug (as defined in paragraph (2) of section 1834(x)) for which the payment amount for a calendar quarter under paragraph (3)(A)(ii)(I) of such section for such quarter exceeds the inflation-adjusted payment under paragraph (3)(A)(ii)(II) of such section for such quarter, the amounts paid shall be the difference between (i) the payment amount under paragraph (3)(A)(ii)(I) of such section for such drug, and (ii) 20 percent of the inflation-adjusted payment
amount under paragraph (3)(A)(ii)(II) of such section for such drug’’;

(B) by adding at the end of the flush left matter following paragraph (9), the following:

“For purposes of applying paragraph (1)(DD), subsections (i)(9) and (t)(3)(H), and section 1834(x)(5), the Secretary shall make such estimates and use such data as the Secretary determines appropriate, and notwithstanding any other provision of law, may do so by program instruction or otherwise.”;

(2) in subsection (i), by adding at the end the following new paragraph:

“(9) In the case of a part B rebatable drug (as defined in paragraph (2) of section 1834(x)) furnished on or after July 1, 2021, under the system under this subsection, in lieu of calculation of coinsurance and the amount of payment otherwise applicable under this subsection, the provisions of section 1834(x)(5), paragraph (1)(DD) of subsection (a), and the flush left matter following paragraph (9) of subsection (a), shall, as determined appropriate by the Secretary, apply under this subsection in the same manner as such provisions of section 1834(x)(5) and subsection (a) apply under such section and subsection.”; and

(3) in subsection (t)(3), by adding at the end the following new subparagraph:
“(H) PART B REBATABLE DRUGS.—In the case of a part B rebatable drug (as defined in paragraph (2) of section 1834(x)) furnished on or after July 1, 2021, under the system under this subsection, in lieu of calculation of coinsurance and the amount of payment otherwise applicable under this subsection, the provisions of section 1834(x)(5), paragraph (1)(DD) of subsection (a), and the flush left matter following paragraph (9) of subsection (a), shall, as determined appropriate by the Secretary, apply under this subsection in the same manner as such provisions of section 1834(x)(5) and subsection (a) apply under such section and subsection.”.

(c) CONFORMING AMENDMENT TO PART B ASP CALCULATION.—Section 1847A(c)(3) of the Social Security Act (42 U.S.C. 1395w–3a(c)(3)) is amended by inserting “or section 1834(x)” after “section 1927”.

SEC. 202. MEDICARE PART D REBATE BY MANUFACTURERS.

(a) IN GENERAL.—Part D of title XVIII of the Social Security Act is amended by inserting after section 1860D–14A (42 U.S.C. 1395w–114a) the following new section:
“SEC. 1860D–14B. MANUFACTURER REBATE FOR CERTAIN DRUGS WITH PRICES INCREASING FASTER THAN INFLATION.

“(a) In General.—

“(1) In General.—Subject to the provisions of this section, in order for coverage to be available under this part for a part D rebatable drug (as defined in subsection (h)(1)) of a manufacturer (as defined in section 1927(k)(5)) dispensed during an applicable year, the manufacturer must have entered into and have in effect an agreement described in subsection (b).

“(2) Authorizing Coverage for Drugs Not Covered Under Agreements.—Paragraph (1) shall not apply to the dispensing of a covered part D drug if—

“(A) the Secretary has made a determination that the availability of the drug is essential to the health of beneficiaries under this part; or

“(B) the Secretary determines that in the period beginning on January 1, 2022, and ending on December 31, 2022, there were extenuating circumstances.

“(3) Applicable Year.—For purposes of this section the term ‘applicable year’ means a year beginning with 2022.
“(b) AGREEMENTS.—

“(1) TERMS OF AGREEMENT.—An agreement described in this subsection, with respect to a manufacturer of a part D rebatable drug, is an agreement under which the following shall apply:

“(A) SECRETARIAL PROVISION OF INFORMATION.—Not later than 9 months after the end of each applicable year with respect to which the agreement is in effect, the Secretary, for each part D rebatable drug of the manufacturer, shall report to the manufacturer the following for such year:

“(i) Information on the total number of units (as defined in subsection (h)(2)) for each dosage form and strength with respect to such part D rebatable drug and year.

“(ii) Information on the amount (if any) of the excess average manufacturer price increase described in subsection (c)(1)(B) for each dosage form and strength with respect to such drug and year.

“(iii) The rebate amount specified under subsection (c) for each dosage form and strength with respect to such drug and year.
“(B) MANUFACTURER REQUIREMENTS.—

For each applicable year with respect to which the agreement is in effect, the manufacturer of the part D rebatable drug, for each dosage form and strength with respect to such drug, not later than 30 days after the date of receipt from the Secretary of the information described in subparagraph (A) for such year, shall provide to the Secretary a rebate that is equal to the amount specified in subsection (c) for such dosage form and strength with respect to such drug for such year.

“(2) LENGTH OF AGREEMENT.—

“(A) IN GENERAL.—An agreement under this section, with respect to a part D rebatable drug, shall be effective for an initial period of not less than one year and shall be automatically renewed for a period of not less than one year unless terminated under subparagraph (B).

“(B) TERMINATION.—

“(i) BY SECRETARY.—The Secretary may provide for termination of an agreement under this section for violation of the requirements of the agreement or other good cause shown. Such termination shall not be
effective earlier than 30 days after the date
of notice of such termination. The Secretary
shall provide, upon request, a manufacturer
with a hearing concerning such a termi-
nation, but such hearing shall not delay the
effective date of the termination.

“(ii) By a manufacturer.—A manu-
ufacturer may terminate an agreement under
this section for any reason. Any such termi-
nation shall be effective, with respect to a
plan year—

“(I) if the termination occurs be-
fore January 30 of the plan year, as of
the day after the end of the plan year;
and

“(II) if the termination occurs on
or after January 30 of the plan year,
as of the day after the end of the suc-
ceeding plan year.

“(C) Effectiveness of termination.—
Any termination under this paragraph shall not
affect rebates due under the agreement under this
section before the effective date of its termination.

“(D) Delay before reentry.—In the
case of any agreement under this section with a


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manufacturer that is terminated in a plan year, the Secretary may not enter into another such agreement with the manufacturer (or a successor manufacturer) before the subsequent plan year, unless the Secretary finds good cause for an earlier reinstatement of such an agreement.

“(c) Rebate Amount.—

“(1) In general.—For purposes of this section, the amount specified in this subsection for a dosage form and strength with respect to a part D rebatable drug and applicable year is, subject to subparagraphs (B) and (C) of paragraph (5), the amount equal to the product of—

“(A) the total number of units of such dosage form and strength with respect to such part D rebatable drug and year; and

“(B) the amount (if any) by which—

“(i) the annual manufacturer price (as determined in paragraph (2)) paid for such dosage form and strength with respect to such part D rebatable drug for the year; ex-

“(ii) the inflation-adjusted payment amount determined under paragraph (3) for
such dosage form and strength with respect to such part D rebatable drug for the year.

“(2) DETERMINATION OF ANNUAL MANUFACTURER PRICE.—The annual manufacturer price determined under this paragraph for a dosage form and strength, with respect to a part D rebatable drug and an applicable year, is the sum of the products of—

“(A) the average manufacturer price (as defined in subsection (h)(6)) of such dosage form and strength, as calculated for a unit of such drug, with respect to each of the calendar quarters of such year; and

“(B) the ratio of—

“(i) the total number of units of such dosage form and strength dispensed during each such calendar quarter of such year; to

“(ii) the total number of units of such dosage form and strength dispensed during such year.

“(3) DETERMINATION OF INFLATION-ADJUSTED PAYMENT AMOUNT.—The inflation-adjusted payment amount determined under this paragraph for a dosage form and strength with respect to a part D rebatable drug for an applicable year, subject to subparagraphs (A) and (D) of paragraph (5), is—
“(A) the benchmark year manufacturer price determined under paragraph (4) for such dosage form and strength with respect to such drug and an applicable year; increased by

“(B) the percentage by which the applicable year CPI–U (as defined in subsection (h)(5)) for the applicable year exceeds the benchmark period CPI–U (as defined in subsection (h)(4)).

“(4) DETERMINATION OF BENCHMARK YEAR MANUFACTURER PRICE.—The benchmark year manufacturer price determined under this paragraph for a dosage form and strength, with respect to a part D rebatable drug and an applicable year, is the sum of the products of—

“(A) the average manufacturer price (as defined in subsection (h)(6)) of such dosage form and strength, as calculated for a unit of such drug, with respect to each calendar quarter of the payment amount benchmark year (as defined in subsection (h)(3)); and

“(B) the ratio of—

“(i) the total number of units of such dosage form and strength dispensed during such calendar quarter of the payment amount benchmark year; to
“(ii) the total number of units of such dosage form and strength dispensed during the payment amount benchmark year.

“(5) SPECIAL TREATMENT OF CERTAIN DRUGS AND EXEMPTION.—

“(A) SUBSEQUENTLY APPROVED DRUGS.—

In the case of a part D rebatable drug first approved or licensed by the Food and Drug Administration after January 1, 2016, subparagraphs (A) and (B) of paragraph (4) shall be applied as if the term ‘payment amount benchmark year’ were defined under subsection (h)(3) as the first calendar year beginning after the day on which the drug was first marketed by any manufacturer and subparagraph (B) of paragraph (3) shall be applied as if the term ‘benchmark period CPI–U’ were defined under subsection (h)(4) as if the reference to ‘January 2016’ under such subsection were a reference to ‘January of the first year beginning after the date on which the drug was first marketed by any manufacturer’.

“(B) EXEMPTION FOR SHORTAGES.—The Secretary may reduce or waive the rebate under paragraph (1) with respect to a part D rebatable drug that is described as currently in shortage
on the shortage list in effect under section 506E of the Federal Food, Drug, and Cosmetic Act or in the case of other exigent circumstances, as determined by the Secretary.

“(C) Treatment of New Formulations.—

“(i) In general.—In the case of a part D rebatable drug that is a line extension of a part D rebatable drug that is an oral solid dosage form, the Secretary shall establish a formula for determining the amount specified in this subsection with respect to such part D rebatable drug and an applicable year with consideration of the original part D rebatable drug.

“(ii) Line extension defined.—In this subparagraph, the term ‘line extension’ means, with respect to a part D rebatable drug, a new formulation of the drug (as determined by the Secretary), such as an extended release formulation, but does not include an abuse-deterrent formulation of the drug (as determined by the Secretary), regardless of whether such abuse-deterrent for-
mulation is an extended release formulation.

“(D) SELECTED DRUGS.—In the case of a part D rebatable drug that is a selected drug (as defined in section 1192(c)) for a price applicability period (as defined in section 1191(b)(2)) and is determined (pursuant to such section 1192(c)) to no longer be a selected drug, for each applicable year beginning after the price applicability period with respect to such drug, subparagraphs (A) and (B) of paragraph (4) shall be applied as if the term ‘payment amount benchmark year’ were defined under subsection (h)(3) as the last year beginning during such price applicability period with respect to such selected drug and subparagraph (B) of paragraph (3) shall be applied as if the term ‘benchmark period CPI–U’ were defined under subsection (h)(4) as if the reference to ‘January 2016’ under such subsection were a reference to January of the last year beginning during such price applicability period with respect to such drug.

“(d) REBATE DEPOSITS.—Amounts paid as rebates under subsection (c) shall be deposited into the Medicare
Prescription Drug Account in the Federal Supplementary Medical Insurance Trust Fund established under section 1841.

“(e) INFORMATION.—For purposes of carrying out this section, the Secretary shall use information submitted by manufacturers under section 1927(b)(3).

“(f) CIVIL MONEY PENALTY.—In the case of a manufacturer of a part D rebatable drug with an agreement in effect under this section who has failed to comply with the terms of the agreement under subsection (b)(1)(B) with respect to such drug for an applicable year, the Secretary may impose a civil money penalty on such manufacturer in an amount equal to 125 percent of the amount specified in subsection (c) for such drug for such year. The provisions of section 1128A (other than subsections (a) (with respect to amounts of penalties or additional assessments) and (b)) shall apply to a civil money penalty under this subsection in the same manner as such provisions apply to a penalty or proceeding under section 1128A(a).

“(g) JUDICIAL REVIEW.—There shall be no judicial review of the following:

“(1) The determination of units under this section.

“(2) The determination of whether a drug is a part D rebatable drug under this section.
“(3) The calculation of the rebate amount under this section.

“(h) DEFINITIONS.—In this section:

“(1) PART D REBATABLE DRUG DEFINED.—

“(A) IN GENERAL.—The term ‘part D rebatable drug’ means a drug or biological that would (without application of this section) be a covered part D drug, except such term shall, with respect to an applicable year, not include such a drug or biological if the average annual total cost under this part for such year per individual who uses such a drug or biological, as determined by the Secretary, is less than, subject to subparagraph (B), $100, as determined by the Secretary using the most recent data available or, if data is not available, as estimated by the Secretary.

“(B) INCREASE.—The dollar amount applied under subparagraph (A)—

“(i) for 2023, shall be the dollar amount specified under such subparagraph for 2022, increased by the percentage increase in the consumer price index for all urban consumers (United States city aver-
age) for the 12-month period beginning with January of 2022; and

“(ii) for a subsequent year, shall be the dollar amount specified in this subparagraph for the previous year, increased by the percentage increase in the consumer price index for all urban consumers (United States city average) for the 12-month period beginning with January of the previous year.

Any dollar amount specified under this subparagraph that is not a multiple of $10 shall be rounded to the nearest multiple of $10.

“(2) UNIT DEFINED.—The term ‘unit’ means, with respect to a part D rebatable drug, the lowest identifiable quantity (such as a capsule or tablet, milligram of molecules, or grams) of the part D rebatable drug that is dispensed to individuals under this part.

“(3) PAYMENT AMOUNT BENCHMARK YEAR.—The term ‘payment amount benchmark year’ means the year beginning January 1, 2016.

“(4) BENCHMARK PERIOD CPI–U.—The term ‘benchmark period CPI–U’ means the consumer price index for all urban consumers (United States city average) for January 2016.
“(5) APPLICABLE YEAR CPI–U.—The term ‘ap-
pllicable year CPI–U’ means, with respect to an ap-
pllicable year, the consumer price index for all urban
consumers (United States city average) for January
of such year.

“(6) AVERAGE MANUFACTURER PRICE.—The
term ‘average manufacturer price’ has the meaning,
with respect to a part D rebatable drug of a manufac-
turer, given such term in section 1927(k)(1), with re-
pect to a covered outpatient drug of a manufacturer
for a rebate period under section 1927.”.

(b) CONFORMING AMENDMENT TO PART B ASP CAL-
CULATION.—Section 1847A(c)(3) of the Social Security Act
(42 U.S.C. 1395w–3a(c)(3)), as amended by section 201(c),
is further amended by striking “section 1927 or section
1834(x)” and inserting “section 1927, section 1834(x), or
section 1860D–14B”.

TITLE III—PART D IMPROVE-
MENTS AND MAXIMUM OUT-
OF-POCKET CAP FOR MED-
CARE BENEFICIARIES

SEC. 301. MEDICARE PART D BENEFIT REDESIGN.

(a) BENEFIT STRUCTURE REDESIGN.—Section
1860D–2(b) of the Social Security Act (42 U.S.C. 1395w–
102(b)) is amended—
(1) in paragraph (2)—

(A) in subparagraph (A), in the matter preceding clause (i), by inserting “for a year preceding 2022 and for costs above the annual deductible specified in paragraph (1) and up to the annual out-of-pocket threshold specified in paragraph (4)(B) for 2022 and each subsequent year” after “paragraph (3)”;

(B) in subparagraph (C)—

(i) in clause (i), in the matter preceding subclause (I), by inserting “for a year preceding 2022,” after “paragraph (4),”;

(ii) in clause (ii)(III), by striking “and each subsequent year” and inserting “and 2021”; and

(C) in subparagraph (D)—

(i) in clause (i)—

(I) in the matter preceding subclause (I), by inserting “for a year preceding 2022,” after “paragraph (4),”;

and

(II) in subclause (I)(bb), by striking “a year after 2018” and inserting
“each of years 2018 through 2021”;

and

(ii) in clause (ii)(V), by striking “2019 and each subsequent year” and inserting “each of years 2019 through 2021”;

(2) in paragraph (3)(A)—

(A) in the matter preceding clause (i), by inserting “for a year preceding 2022,” after “and (4),”; and

(B) in clause (ii), by striking “for a subsequent year” and inserting “for each of years 2007 through 2021”; and

(3) in paragraph (4)—

(A) in subparagraph (A)—

(i) in clause (i)—

(II) by redesignating subclauses (I) and (II) as items (aa) and (bb), respectively, and moving the margin of each such redesignated item 2 ems to the right;

(II) in the matter preceding item (aa), as redesignated by subclause (I), by striking “is equal to the greater of—” and inserting “is equal to—
“(I) for a year preceding 2022, the greater of—”;

(III) by striking the period at the end of item (bb), as redesignated by subclause (I), and inserting “; and”;

and

(IV) by adding at the end the following:

“(II) for 2022 and each succeeding year, $0.”; and

(ii) in clause (ii), by striking “clause (i)(I)” and inserting “clause (i)(I)(aa)”;

(B) in subparagraph (B)—

(i) in clause (i)—

(I) in subclause (V), by striking “or” at the end;

(II) in subclause (VI)—

(aa) by striking “for a subsequent year” and inserting “for 2021”; and

(bb) by striking the period at the end and inserting a semicolon;

and

(III) by adding at the end the following new subclauses:
“(VII) for 2022, is equal to $2,000; or
“(VIII) for a subsequent year, is equal to the amount specified in this subparagraph for the previous year, increased by the annual percentage increase described in paragraph (6) for the year involved.”; and

(ii) in clause (ii), by striking “clause (i)(II)” and inserting “clause (i)”;

(C) in subparagraph (C)(i), by striking “and for amounts” and inserting “and, for a year preceding 2022, for amounts”; and

(D) in subparagraph (E), by striking “In applying” and inserting “For each of years 2011 through 2021, in applying”.

(b) DECREASING REINSURANCE PAYMENT AMOUNT.—

Section 1860D–15(b)(1) of the Social Security Act (42 U.S.C. 1395w–115(b)(1)) is amended by inserting after “80 percent” the following: “(or, with respect to a coverage year after 2021, 20 percent)”.

(c) MANUFACTURER DISCOUNT PROGRAM.—

(1) IN GENERAL.—Part D of title XVIII of the Social Security Act (42 U.S.C. 1395w–101 et seq.), as amended by section 202, is further amended by insert-
ing after section 1860D–14B the following new section:

“SEC. 1860D–14C. MANUFACTURER DISCOUNT PROGRAM.

“(a) Establishment.—The Secretary shall establish a manufacturer discount program (in this section referred to as the ‘program’). Under the program, the Secretary shall enter into agreements described in subsection (b) with manufacturers and provide for the performance of the duties described in subsection (c). The Secretary shall establish a model agreement for use under the program by not later than January 1, 2021, in consultation with manufacturers, and allow for comment on such model agreement.

“(b) Terms of Agreement.—

“(1) In general.—

“(A) Agreement.—An agreement under this section shall require the manufacturer to provide applicable beneficiaries access to discounted prices for applicable drugs of the manufacturer that are dispensed on or after January 1, 2022.

“(B) Provision of discounted prices at the point-of-sale.—The discounted prices described in subparagraph (A) shall be provided to the applicable beneficiary at the pharmacy or by
the mail order service at the point-of-sale of an applicable drug.

“(C) TIMING OF AGREEMENT.—

“(i) SPECIAL RULE FOR 2022.—In order for an agreement with a manufacturer to be in effect under this section with respect to the period beginning on January 1, 2022, and ending on December 31, 2022, the manufacturer shall enter into such agreement not later than 30 days after the date of the establishment of a model agreement under subsection (a).

“(ii) 2023 AND SUBSEQUENT YEARS.—In order for an agreement with a manufacturer to be in effect under this section with respect to plan year 2023 or a subsequent plan year, the manufacturer shall enter into such agreement (or such agreement shall be renewed under paragraph (4)(A)) not later than January 30 of the preceding year.

“(2) PROVISION OF APPROPRIATE DATA.—Each manufacturer with an agreement in effect under this section shall collect and have available appropriate data, as determined by the Secretary, to ensure that
it can demonstrate to the Secretary compliance with the requirements under the program.

“(3) COMPLIANCE WITH REQUIREMENTS FOR ADMINISTRATION OF PROGRAM.—Each manufacturer with an agreement in effect under this section shall comply with requirements imposed by the Secretary or a third party with a contract under subsection (d)(3), as applicable, for purposes of administering the program, including any determination under subparagraph (A) of subsection (c)(1) or procedures established under such subsection (c)(1).

“(4) LENGTH OF AGREEMENT.—

“(A) IN GENERAL.—An agreement under this section shall be effective for an initial period of not less than 12 months and shall be automatically renewed for a period of not less than 1 year unless terminated under subparagraph (B).

“(B) TERMINATION.—

“(i) BY THE SECRETARY.—The Secretary may provide for termination of an agreement under this section for a knowing and willful violation of the requirements of the agreement or other good cause shown. Such termination shall not be effective ear-
lier than 30 days after the date of notice to
the manufacturer of such termination. The
Secretary shall provide, upon request, a
manufacturer with a hearing concerning
such a termination, and such hearing shall
take place prior to the effective date of the
termination with sufficient time for such ef-
tective date to be repealed if the Secretary
determines appropriate.

“(ii) BY A MANUFACTURER.—A manu-
facturer may terminate an agreement under
this section for any reason. Any such termi-
nation shall be effective, with respect to a
plan year—

“(I) if the termination occurs be-
fore January 30 of a plan year, as of
the day after the end of the plan year;
and

“(II) if the termination occurs on
or after January 30 of a plan year, as
of the day after the end of the suc-
ceeding plan year.

“(iii) EFFECTIVENESS OF TERMI-
nATION.—Any termination under this sub-
paragraph shall not affect discounts for ap-
licable drugs of the manufacturer that are due under the agreement before the effective date of its termination.

“(iv) NOTICE TO THIRD PARTY.—The Secretary shall provide notice of such termination to a third party with a contract under subsection (d)(3) within not less than 30 days before the effective date of such termination.

“(c) DUTIES DESCRIBED.—The duties described in this subsection are the following:

“(1) ADMINISTRATION OF PROGRAM.—Administering the program, including—

“(A) the determination of the amount of the discounted price of an applicable drug of a manufacturer;

“(B) the establishment of procedures under which discounted prices are provided to applicable beneficiaries at pharmacies or by mail order service at the point-of-sale of an applicable drug;

“(C) the establishment of procedures to ensure that, not later than the applicable number of calendar days after the dispensing of an applicable drug by a pharmacy or mail order service, the pharmacy or mail order service is reim-
bursed for an amount equal to the difference be-
 tween—

“(i) the negotiated price of the applica-
 ble drug; and

“(ii) the discounted price of the appli-
 cable drug;

“(D) the establishment of procedures to en-
 sure that the discounted price for an applicable
 drug under this section is applied before any
 coverage or financial assistance under other
 health benefit plans or programs that provide
 coverage or financial assistance for the purchase
 or provision of prescription drug coverage on be-
 half of applicable beneficiaries as the Secretary
 may specify; and

“(E) providing a reasonable dispute resolu-
 tion mechanism to resolve disagreements between
 manufacturers, applicable beneficiaries, and the
 third party with a contract under subsection
 (d)(3).

“(2) MONITORING COMPLIANCE.—

“(A) In general.—The Secretary shall
 monitor compliance by a manufacturer with the
terms of an agreement under this section.
“(B) Notification.—If a third party with a contract under subsection (d)(3) determines that the manufacturer is not in compliance with such agreement, the third party shall notify the Secretary of such noncompliance for appropriate enforcement under subsection (e).

“(3) Collection of data from prescription drug plans and MA–PD plans.—The Secretary may collect appropriate data from prescription drug plans and MA–PD plans in a timeframe that allows for discounted prices to be provided for applicable drugs under this section.

“(d) Administration.—

“(1) In general.—Subject to paragraph (2), the Secretary shall provide for the implementation of this section, including the performance of the duties described in subsection (c).

“(2) Limitation.—In providing for the implementation of this section, the Secretary shall not receive or distribute any funds of a manufacturer under the program.

“(3) Contract with third parties.—The Secretary shall enter into a contract with 1 or more third parties to administer the requirements established by the Secretary in order to carry out this sec-
tion. At a minimum, the contract with a third party under the preceding sentence shall require that the third party—

“(A) receive and transmit information between the Secretary, manufacturers, and other individuals or entities the Secretary determines appropriate;

“(B) receive, distribute, or facilitate the distribution of funds of manufacturers to appropriate individuals or entities in order to meet the obligations of manufacturers under agreements under this section;

“(C) provide adequate and timely information to manufacturers, consistent with the agreement with the manufacturer under this section, as necessary for the manufacturer to fulfill its obligations under this section; and

“(D) permit manufacturers to conduct periodic audits, directly or through contracts, of the data and information used by the third party to determine discounts for applicable drugs of the manufacturer under the program.

“(4) PERFORMANCE REQUIREMENTS.—The Secretary shall establish performance requirements for a third party with a contract under paragraph (3) and
safeguards to protect the independence and integrity
of the activities carried out by the third party under
the program under this section.

“(5) **IMPLEMENTATION.**—Notwithstanding any
other provision of law, the Secretary may implement
the program under this section by program instruc-
tion or otherwise.

“(6) **ADMINISTRATION.**—Chapter 35 of title 44,
United States Code, shall not apply to the program
under this section.

“(e) **ENFORCEMENT.**—

“(1) **AUDITS.**—Each manufacturer with an
agreement in effect under this section shall be subject
to periodic audit by the Secretary.

“(2) **CIVIL MONEY PENALTY.**—

“(A) **IN GENERAL.**—The Secretary may im-
pose a civil money penalty on a manufacturer
that fails to provide applicable beneficiaries dis-
counts for applicable drugs of the manufacturer
in accordance with such agreement for each such
failure in an amount the Secretary determines is
equal to the sum of—

“(i) the amount that the manufacturer
would have paid with respect to such dis-
counts under the agreement, which will then
be used to pay the discounts which the manufacturer had failed to provide; and

“(ii) 25 percent of such amount.

“(B) APPLICATION.—The provisions of section 1128A (other than subsections (a) and (b)) shall apply to a civil money penalty under this paragraph in the same manner as such provisions apply to a penalty or proceeding under section 1128A(a).

“(f) CLARIFICATION REGARDING AVAILABILITY OF OTHER COVERED PART D DRUGS.—Nothing in this section shall prevent an applicable beneficiary from purchasing a covered part D drug that is not an applicable drug (including a generic drug or a drug that is not on the formulary of the prescription drug plan or MA–PD plan that the applicable beneficiary is enrolled in).

“(g) DEFINITIONS.—In this section:

“(1) APPLICABLE BENEFICIARY.—The term ‘applicable beneficiary’ means an individual who, on the date of dispensing a covered part D drug—

“(A) is enrolled in a prescription drug plan or an MA–PD plan;

“(B) is not enrolled in a qualified retiree prescription drug plan; and
“(C) has incurred costs for covered part D drugs in the year that are equal to or exceed the annual deductible specified in section 1860D–2(b)(1) for such year.

“(2) APPLICABLE DRUG.—The term ‘applicable drug’, with respect to an applicable beneficiary—

“(A) means a covered part D drug—

“(i) approved under a new drug application under section 505(c) of the Federal Food, Drug, and Cosmetic Act or, in the case of a biologic product, licensed under section 351 of the Public Health Service Act; and

“(ii)(I) if the PDP sponsor of the prescription drug plan or the MA organization offering the MA–PD plan uses a formulary, which is on the formulary of the prescription drug plan or MA–PD plan that the applicable beneficiary is enrolled in;

“(II) if the PDP sponsor of the prescription drug plan or the MA organization offering the MA–PD plan does not use a formulary, for which benefits are available under the prescription drug plan or MA–
PD plan that the applicable beneficiary is enrolled in; or

“(III) is provided through an exception or appeal; and

“(B) does not include a selected drug (as defined in section 1192(c)) during a price applicability period (as defined in section 1191(b)(2)) with respect to such drug.

“(3) APPLICABLE NUMBER OF CALENDAR DAYS.—The term ‘applicable number of calendar days’ means—

“(A) with respect to claims for reimbursement submitted electronically, 14 days; and

“(B) with respect to claims for reimbursement submitted otherwise, 30 days.

“(4) DISCOUNTED PRICE.—

“(A) IN GENERAL.—The term ‘discounted price’ means, with respect to an applicable drug of a manufacturer furnished during a year to an applicable beneficiary—

“(i) who has not incurred costs for covered part D drugs in the year that are equal to or exceed the annual out-of-pocket threshold specified in section 1860D—
2(b)(4)(B)(i) for the year, 90 percent of the negotiated price of such drug; and

“(ii) who has incurred such costs in the year that are equal to or exceed such threshold for the year, 70 percent of the negotiated price of such drug.

“(B) CLARIFICATION.—Nothing in this section shall be construed as affecting the responsibility of an applicable beneficiary for payment of a dispensing fee for an applicable drug.

“(C) SPECIAL CASE FOR CERTAIN CLAIMS.—

“(i) CLAIMS SPANNING DEDUCTIBLE.—

In the case where the entire amount of the negotiated price of an individual claim for an applicable drug with respect to an applicable beneficiary does not fall at or above the annual deductible specified in section 1860D–2(b)(1) for the year, the manufacturer of the applicable drug shall provide the discounted price under this section on only the portion of the negotiated price of the applicable drug that falls at or above such annual deductible.
“(ii) Claims spanning out-of-pocket threshold.—In the case where the entire amount of the negotiated price of an individual claim for an applicable drug with respect to an applicable beneficiary does not fall entirely below or entirely above the annual out-of-pocket threshold specified in section 1860D–2(b)(4)(B)(i) for the year, the manufacturer of the applicable drug shall provide the discounted price—

“(I) in accordance with subparagraph (A)(i) on the portion of the negotiated price of the applicable drug that falls below such threshold; and

“(II) in accordance with subparagraph (A)(ii) on the portion of such price of such drug that falls at or above such threshold.

“(5) Manufacturer.—The term ‘manufacturer’ means any entity which is engaged in the production, preparation, propagation, compounding, conversion, or processing of prescription drug products, either directly or indirectly by extraction from substances of natural origin, or independently by means of chemical synthesis, or by a combination of extraction and
chemical synthesis. Such term does not include a wholesale distributor of drugs or a retail pharmacy licensed under State law.

“(6) NEGOTIATED PRICE.—The term ‘negotiated price’ has the meaning given such term in section 423.100 of title 42, Code of Federal Regulations (or any successor regulation), except that, with respect to an applicable drug, such negotiated price shall not include any dispensing fee for the applicable drug.

“(7) QUALIFIED RETIREE PRESCRIPTION DRUG PLAN.—The term ‘qualified retiree prescription drug plan’ has the meaning given such term in section 1860D–22(a)(2).”.

(2) SUNSET OF MEDICARE COVERAGE GAP DISCOUNT PROGRAM.—Section 1860D–14A of the Social Security Act (42 U.S.C. 1395–114a) is amended—

(A) in subsection (a), in the first sentence, by striking “The Secretary” and inserting “Subject to subsection (h), the Secretary”;

(B) by adding at the end the following new subsection:

“(h) SUNSET OF PROGRAM.—

“(1) IN GENERAL.—The program shall not apply with respect to applicable drugs dispensed on or after January 1, 2022, and, subject to paragraph (2),
agreements under this section shall be terminated as of such date.

“(2) CONTINUED APPLICATION FOR APPLICABLE DRUGS DISPENSED PRIOR TO SUNSET.—The provisions of this section (including all responsibilities and duties) shall continue to apply after January 1, 2022, with respect to applicable drugs dispensed prior to such date.”.

(3) INCLUSION OF ACTUARIAL VALUE OF MANUFACTURER DISCOUNTS IN BIDS.—Section 1860D–11 of the Social Security Act (42 U.S.C. 1395w–111) is amended—

(A) in subsection (b)(2)(C)(iii)—

(i) by striking “assumptions regarding the reinsurance” and inserting “assumptions regarding—

“(I) the reinsurance”; and

(ii) by adding at the end the following:

“(II) for 2022 and each subsequent year, the manufacturer discounts provided under section 1860D–14C subtracted from the actuarial value to produce such bid; and”; and

(B) in subsection (c)(1)(C)—
(i) by striking “an actuarial valuation of the reinsurance” and inserting “an actuarial valuation of—

“(i) the reinsurance”;

(ii) in clause (i), as inserted by clause (i) of this subparagraph, by adding “and” at the end; and

(iii) by adding at the end the following:

“(ii) for 2022 and each subsequent year, the manufacturer discounts provided under section 1860D–14C;”.

(d) CONFORMING AMENDMENTS.—

(1) Section 1860D–2 of the Social Security Act (42 U.S.C. 1395w–102) is amended—

(A) in subsection (a)(2)(A)(i)(I), by striking “, or an increase in the initial” and inserting “or, for a year preceding 2022, an increase in the initial”;

(B) in subsection (c)(1)(C)—

(i) in the subparagraph heading, by striking “AT INITIAL COVERAGE LIMIT”; and

(ii) by inserting “for a year preceding 2022 or the annual out-of-pocket threshold specified in subsection (b)(4)(B) for the year
for 2022 and each subsequent year” after “subsection (b)(3) for the year” each place it appears; and
(C) in subsection (d)(1)(A), by striking “or an initial” and inserting “or, for a year preceding 2022, an initial”.

(2) Section 1860D–4(a)(4)(B)(i) of the Social Security Act (42 U.S.C. 1395w–104(a)(4)(B)(i)) is amended by striking “the initial” and inserting “for a year preceding 2022, the initial”.

(3) Section 1860D–14(a) of the Social Security Act (42 U.S.C. 1395w–114(a)) is amended—
(A) in paragraph (1)—
(i) in subparagraph (C), by striking “The continuation” and inserting “For a year preceding 2022, the continuation”;
(iii) in subparagraph (E), by striking “The elimination” and inserting “For a year preceding 2022, the elimination”; and
(B) in paragraph (2)—
(i) in subparagraph (C), by striking “The continuation” and inserting “For a year preceding 2022, the continuation”; and


(A) by striking “the value of any discount” and inserting the following: “the value of—

“(i) for years prior to 2022, any discount”; 

(B) in clause (i), as inserted by subparagraph (A) of this paragraph, by striking the period at the end and inserting “; and”; and 

(C) by adding at the end the following new clause:

“(ii) for 2022 and each subsequent year, any discount provided pursuant to section 1860D–14C.”.
(6) Section 1860D–41(a)(6) of the Social Security Act (42 U.S.C. 1395w–151(a)(6)) is amended—

(A) by inserting “for a year before 2022” after “1860D–2(b)(3)”; and

(B) by inserting “for such year” before the period.

(7) Section 1860D–43 of the Social Security Act (42 U.S.C. 1395w–153) is amended—

(A) in subsection (a)—

(i) by striking paragraph (1) and inserting the following:

“(1) participate in—

“(A) for 2011 through 2021, the Medicare coverage gap discount program under section 1860D–14A; and

“(B) for 2022 and each subsequent year, the manufacturer discount program under section 1860D–14C;”;

(ii) by striking paragraph (2) and inserting the following:

“(2) have entered into and have in effect—

“(A) for 2011 through 2021, an agreement described in subsection (b) of section 1860D–14A with the Secretary; and
“(B) for 2022 and each subsequent year, an
agreement described in subsection (b) of section
1860D–14C with the Secretary; and”; and
(iii) by striking paragraph (3) and insert-
sing the following:
“(3) have entered into and have in effect, under
terms and conditions specified by the Secretary—
“(A) for 2011 through 2021, a contract with
a third party that the Secretary has entered into
a contract with under subsection (d)(3) of section
1860D–14A; and
“(B) for 2022 and each subsequent year, a
contract with a third party that the Secretary
has entered into a contract with under subsection
(d)(3) of section 1860D–14C.”; and
(B) by striking subsection (b) and inserting
the following:
“(b) EFFECTIVE DATE.—Paragraphs (1)(A), (2)(A),
and (3)(A) of subsection (a) shall apply to covered part D
drugs dispensed under this part on or after January 1,
2011, and before January 1, 2022, and paragraphs (1)(B),
(2)(B), and (3)(B) of such subsection shall apply to covered
part D drugs dispensed under this part on or after January
1, 2022.”.
(e) EFFECTIVE DATE.—The amendments made by this section shall apply with respect to plan year 2022 and subsequent plan years.

SEC. 302. ALLOWING CERTAIN ENROLLEES OF PRESCRIPTION DRUGS PLANS AND MA–PD PLANS UNDER MEDICARE PROGRAM TO SPREAD OUT COST-SHARING UNDER CERTAIN CIRCUMSTANCES.

Section 1860D–2(b)(2) of the Social Security Act (42 U.S.C. 1395w–102(b)(2)), as amended by section 301, is further amended—

(1) in subparagraph (A), by striking “Subject to subparagraphs (C) and (D)” and inserting “Subject to subparagraphs (C), (D), and (E)”;

(2) by adding at the end the following new subparagraph:

“(E) ENROLLEE OPTION REGARDING SPREADING COST-SHARING.—The Secretary shall establish by regulation a process under which, with respect to plan year 2022 and subsequent plan years, a prescription drug plan or an MA–PD plan shall, in the case of a part D eligible individual enrolled with such plan for such plan year who is not a subsidy eligible individual (as defined in section 1860D–14(a)(3)) and with re-
spect to whom the plan projects that the dispensing of the first fill of a covered part D drug to such individual will result in the individual incurring costs that are equal to or above the annual out-of-pocket threshold specified in paragraph (4)(B) for such plan year, provide such individual with the option to make the coinsurance payment required under subparagraph (A) (for the portion of such costs that are not above such annual out-of-pocket threshold) in the form of periodic installments over the remainder of such plan year.”.

SEC. 303. ESTABLISHMENT OF PHARMACY QUALITY MEASURES UNDER MEDICARE PART D.

Section 1860D–4(c) of the Social Security Act (42 U.S.C. 1395w–104(c)) is amended—

(1) by redesignating the paragraph (6), as added by section 50354 of division E of the Bipartisan Budget Act of 2018 (Public Law 115–123), as paragraph (7); and

(2) by adding at the end the following new paragraph:

“(8) APPLICATION OF PHARMACY QUALITY MEASURES.—
“(A) IN GENERAL.—A PDP sponsor that implements incentive payments to a pharmacy or price concessions paid by a pharmacy based on quality measures shall use measures established or approved by the Secretary under subparagraph (B) with respect to payment for covered part D drugs dispensed by such pharmacy.

“(B) STANDARD PHARMACY QUALITY MEASURES.—The Secretary shall establish or approve standard quality measures from a consensus and evidence-based organization for payments described in subparagraph (A). Such measures shall focus on patient health outcomes and be based on proven criteria measuring pharmacy performance.

“(C) EFFECTIVE DATE.—The requirement under subparagraph (A) shall take effect for plan years beginning on or after January 1, 2021, or such earlier date specified by the Secretary if the Secretary determines there are sufficient measures established or approved under subparagraph (B) to meet the requirement under subparagraph (A).”
TITLE IV—PRESCRIPTION DRUG POLICIES FOR LOW-INCOME INDIVIDUALS

SEC. 401. ADJUSTMENTS TO MEDICARE PART D COST-SHARING REDUCTIONS FOR LOW-INCOME INDIVIDUALS.

Section 1860D–14(a) of the Social Security Act (42 U.S.C. 1395w–114(a)), as amended by section 301(d), is further amended—

(1) in paragraph (1)—

(A) in subparagraph (D)—

(i) in clause (ii)—

(I) by striking “that does not exceed $1 for” and all that follows through the period at the end and inserting “that does not exceed—

“(I) for plan years before plan year 2021—

“(aa) for a generic drug or a preferred drug that is a multiple source drug (as defined in section 1927(k)(7)(A)(i)), $1 or, if less, the copayment amount applicable to an individual under clause (iii); and
“(bb) for any other drug, $3
or, if less, the copayment amount
applicable to an individual under
clause (iii); and”; and

(II) by adding at the end the fol-
lowing new subclauses:

“(II) for plan year 2021—

“(aa) for a generic drug, $0;
and

“(bb) for any other drug, the
dollar amount applied under this
clause (after application of para-
graph (4)(A)) for plan year 2020
for a drug described in subclause
(I)(bb); and

“(III) for a subsequent year, the
dollar amount applied under this
clause for the previous year for the
drug, increased by the annual percent-
age increase in the consumer price
index (all items; U.S. city average) as
of September of such previous year.”;
and

(ii) in clause (iii)—
(I) by striking “does not exceed the copayment amount specified under” and inserting “does not exceed—

“(I) for plan years beginning before plan year 2021, the copayment amount specified under”;

(II) by striking the period at the end and inserting “; and”;

(III) by adding at the end the following new subclause:

“(II) for plan year 2021 and each subsequent plan year, the copayment amount applied under clause (ii) for the drug and year involved.”; and

(B) by adding at the end the following new subparagraph:

“(F) ROUNDING.—Any amount established under clause (ii) of subparagraph (D), including as applied under clause (iii) of such subparagraph or paragraph (2)(D), that is based on an increase of $3, that is not a multiple of 5 cents or 10 cents, respectively, shall be rounded to the nearest multiple of 5 cents or 10 cents, respectively.”;
(2) in paragraph (2)—

(A) in subparagraph (D)—

(i) by striking “of coinsurance of” and inserting “of—

“(i) for plan years before plan year 2021, coinsurance of”;

(ii) by striking the period at the end and inserting “; and”; and

(iii) by adding at the end the following new clause:

“(ii) for plan year 2021 and each subsequent plan year, a copayment amount that does not exceed the copayment amount applied under paragraph (1)(D)(ii) for the drug and year involved.”; and

(B) in subparagraph (E)—

(i) by striking “subsection (c), the substitution for” and inserting “subsection (c)—

“(i) for plan years before plan year 2021, the substitution for”; 

(ii) by striking the period at the end and inserting “; and”; and

(iii) by adding at the end the following new clause:
“(ii) for plan year 2021, the elimination of any cost-sharing imposed under section 1860D–2(b)(4)(A).”; and

(3) in paragraph (4)(A)(ii), by inserting “(before 2021)” after “subsequent year”.

SEC. 402. DISSEMINATION TO MEDICARE PART D SUBSIDY ELIGIBLE INDIVIDUALS OF INFORMATION COMPARING PREMIUMS OF CERTAIN PRESCRIPTION DRUG PLANS.

Section 1860D–1(c)(3) of the Social Security Act (42 U.S.C. 1395w–101(c)(3)) is amended by adding at the end the following new subparagraph:

“(C) INFORMATION ON PREMIUMS FOR SUBSIDY ELIGIBLE INDIVIDUALS.—

“(i) IN GENERAL.—For plan year 2022 and each subsequent plan year, the Secretary shall disseminate to each subsidy eligible individual (as defined in section 1860D–14(a)(3)) information under this paragraph comparing premiums that would apply to such individual for prescription drug coverage under LIS benchmark plans, including, in the case of an individual enrolled in a prescription drug plan under this part, information that compares the
premium that would apply if such individual were to remain enrolled in such plan to premiums that would apply if the individual were to enroll in other LIS benchmark plans.

“(ii) LIS BENCHMARK PLAN.—For purposes of clause (i), the term ‘LIS benchmark plan’ means, with respect to an individual, a prescription drug plan under this part that is offered in the region in which the individual resides and—

“(I) that provides for a premium that is not more than the low-income benchmark premium amount (as defined in section 1860D–14(b)(2)) for such region; or

“(II) with respect to which the premium would be waived as de minimis pursuant to section 1860D–14(a)(5) for such individual.”.
SEC. 403. PROVIDING FOR INTELLIGENT ASSIGNMENT OF CERTAIN SUBSIDY ELIGIBLE INDIVIDUALS AUTO-ENROLLED UNDER MEDICARE PRESCRIPTION DRUG PLANS AND MA–PD PLANS.

(a) In General.—Section 1860D–1(b)(1) of the Social Security Act (42 U.S.C. 1395w–101(b)(1)) is amended—

(1) in subparagraph (C)—

(A) by inserting after “PDP region” the following: “or through use of an intelligent assignment process that is designed to maximize the access of such individual to necessary prescription drugs while minimizing costs to such individual and to the program under this part to the greatest extent possible. In the case the Secretary enrolls such individuals through use of an intelligent assignment process, such process shall take into account the extent to which prescription drugs necessary for the individual are covered in the case of a PDP sponsor of a prescription drug plan that uses a formulary, the use of prior authorization or other restrictions on access to coverage of such prescription drugs by such a sponsor, and the overall quality of a prescription drug plan as measured by quality ratings established by the Secretary”; and

...
(B) by striking “Nothing in the previous sentence” and inserting “Nothing in this subparagraph”; and

(2) in subparagraph (D)—

(A) by inserting after “PDP region” the following: “or through use of an intelligent assignment process that is designed to maximize the access of such individual to necessary prescription drugs while minimizing costs to such individual and to the program under this part to the greatest extent possible. In the case the Secretary enrolls such individuals through use of an intelligent assignment process, such process shall take into account the extent to which prescription drugs necessary for the individual are covered in the case of a PDP sponsor of a prescription drug plan that uses a formulary, the use of prior authorization or other restrictions on access to coverage of such prescription drugs by such a sponsor, and the overall quality of a prescription drug plan as measured by quality ratings established by the Secretary”; and

(B) by striking “Nothing in the previous sentence” and inserting “Nothing in this subparagraph”.
(b) **Effective Date.**—The amendments made by subsection (a) shall apply with respect to plan years beginning with plan year 2022.

**Sec. 404. Expanding Eligibility for Low-Income Subsidies Under Part D of the Medicare Program.**

Section 1860D–14(a) of the Social Security Act (42 U.S.C. 1395w–114(a)), as amended by sections 301(d) and 401, is further amended—

(1) in the subsection heading, by striking “INDIVIDUALS” and all that follows through “LINE” and inserting “CERTAIN INDIVIDUALS”;

(2) in paragraph (1)—

(A) by striking the paragraph heading and inserting “INDIVIDUALS WITH CERTAIN LOW INCOMES”; and

(B) in the matter preceding subparagraph (A), by inserting “(or, with respect to a plan year beginning on or after January 1, 2022, 150 percent)” after “135 percent”;

(3) in paragraph (2)—

(A) by striking the paragraph heading and inserting “OTHER LOW-INCOME INDIVIDUALS”; and

(B) in subparagraph (A)—
(i) by inserting “(or, with respect to a plan year beginning on or after January 1, 2022, 150 percent)” after “135 percent”; and

(ii) by inserting “(or, with respect to a plan year beginning on or after January 1, 2022, 200 percent)” after “150 percent”; and

(4) in paragraph (3)(A)(ii), by inserting “(or, with respect to a plan year beginning on or after January 1, 2022, 200 percent)” after “150 percent”.

SEC. 405. AUTOMATIC ELIGIBILITY OF CERTAIN LOW-INCOME TERRITORIAL RESIDENTS FOR PREMIUM AND COST-SHARING SUBSIDIES UNDER THE MEDICARE PROGRAM; SUNSET OF ENHANCED ALLOTMENT PROGRAM.

(a) AUTOMATIC ELIGIBILITY OF CERTAIN LOW-INCOME TERRITORIAL RESIDENTS FOR PREMIUM AND COST-SHARING SUBSIDIES UNDER THE MEDICARE PROGRAM.—

(1) IN GENERAL.—Section 1860D–14(a)(3) of the Social Security Act (42 U.S.C. 1395w–114(a)(3)) is amended—

(A) in subparagraph (B)(v)—

(i) in subclause (I), by striking “and” at the end;
(ii) in subclause (II), by striking the period and inserting "; and"; and

(iii) by inserting after subclause (II) the following new subclause:

"(III) with respect to plan years beginning on or after January 1, 2021, shall provide that any part D eligible individual who is enrolled for medical assistance under the State Medicaid plan of a territory (as defined in section 1935(f)) under title XIX (or a waiver of such a plan) shall be treated as a subsidy eligible individual described in paragraph (1).”; and

(B) in subparagraph (F), by adding at the end the following new sentence: "The previous sentence shall not apply with respect to eligibility determinations for premium and cost-sharing subsidies under this section made on or after January 1, 2021.”.

(2) CONFORMING AMENDMENT.—Section 1860D–31(j)(2)(D) of the Social Security Act (42 U.S.C. 1395w–141(j)(2)(D)) is amended by adding at the end the following new sentence: “The previous sentence shall not apply with respect to amounts made
available to a State under this paragraph on or after January 1, 2021.”

(b) **Sunset of Enhanced Allotment Program.**—

(1) In general.—Section 1935(e) of the Social Security Act (42 U.S.C. 1396u–5(e)) is amended—

(A) in paragraph (1)(A), by inserting after “such State” the following: “before January 1, 2021”; and

(B) in paragraph (3)—

(i) in subparagraph (A), in the matter preceding clause (i), by inserting after “a year” the following: “(before 2021)”;

(ii) in subparagraph (B)(iii), by striking “a subsequent year” and inserting “each of fiscal years 2008 through 2020”.

(2) Territory Defined.—Section 1935 of the Social Security Act (42 U.S.C. 1396u–5) is amended by adding at the end the following new subsection:

“(f) **Territory Defined.**—In this section, the term ‘territory’ means Puerto Rico, the Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa.”.
SEC. 406. AUTOMATIC QUALIFICATION OF CERTAIN MEDICAID BENEFICIARIES FOR PREMIUM AND COST-SHARING SUBSIDIES UNDER PART D OF THE MEDICARE PROGRAM.

Clause (v) of section 1860D–14(a)(3)(B) of the Social Security Act (42 U.S.C. 1395w–114(a)(3)(B)), as amended by section 405, is further amended—

(1) in subclause (II), by striking “and” at the end;

(2) in subclause (III), by striking the period and inserting “; and”;

(3) by inserting after subclause (III) the following new subclause:

“(IV) with respect to plan years beginning on or after January 1, 2022, shall, notwithstanding the preceding clauses of this subparagraph, provide that any part D eligible individual not described in subclause (I), (II), or (III) who is enrolled, as of the day before the date on which such individual attains the age of 65, for medical assistance under a State plan under title XIX (or a waiver of such plan) pursuant to clause (i)(VIII) or (ii)(XX) of section 1902(a)(10)(A), and who has income
below 200 percent of the poverty line applicable to a family of the size involved, shall be treated as a subsidy eligible individual described in paragraph (1) for a limited period of time, as specified by the Secretary.”.

SEC. 407. ELIMINATING THE RESOURCE REQUIREMENT WITH RESPECT TO SUBSIDY ELIGIBLE INDIVIDUALS UNDER PART D OF THE MEDICARE PROGRAM.


SEC. 408. PROVIDING FOR CERTAIN RULES REGARDING THE TREATMENT OF ELIGIBLE RETIREMENT PLANS IN DETERMINING THE ELIGIBILITY OF INDIVIDUALS FOR PREMIUM AND COST-SHARING SUBSIDIES UNDER PART D OF THE MEDICARE PROGRAM.

Section 1860D–14(a)(3)(C)(i) of the Social Security Act (42 U.S.C. 1395w–114(a)(3)(C)(i)) is amended, by striking “except that support and maintenance furnished in kind shall not be counted as income; and” and inserting “except that—
“(I) support and maintenance furnished in kind shall not be counted as income; and
“(II) for plan years beginning on or after January 1, 2022, any distribution or withdrawal from an eligible retirement plan (as defined in subparagraph (B) of section 402(c)(8) of the Internal Revenue Code of 1986, but excluding any defined benefit plan described in clause (iv) or (v) of such subparagraph and any qualified trust (as defined in subparagraph (A) of such section) which is part of such a defined benefit plan) shall be counted as income; and”.

**TITLE V—DRUG PRICE TRANSPARENCY**

**SEC. 501. DRUG PRICE TRANSPARENCY.**

Part A of title XI of the Social Security Act is amended by adding at the end the following new sections:

“SEC. 1150C. REPORTING ON DRUG PRICES.

“(a) DEFINITIONS.—In this section:

“(1) MANUFACTURER.—The term ‘manufacturer’ means the person—
“(A) that holds the application for a drug approved under section 505 of the Federal Food, Drug, and Cosmetic Act or licensed under section 351 of the Public Health Service Act; or

“(B) who is responsible for setting the wholesale acquisition cost for the drug.

“(2) QUALIFYING DRUG.—The term ‘qualifying drug’ means any drug that is approved under subsection (c) or (j) of section 505 of the Federal Food, Drug, and Cosmetic Act or licensed under subsection (a) or (k) of section 351 of the Public Health Service Act—

“(A) that has a wholesale acquisition cost of $100 or more, adjusted for inflation occurring after the date of enactment of this section, for a month’s supply or a typical course of treatment that lasts less than a month, and is—

“(i) subject to section 503(b)(1) of the Federal Food, Drug, and Cosmetic Act; and

“(ii) not a preventative vaccine; and

“(B) for which, during the previous calendar year, at least 1 dollar of the total amount of sales were for individuals enrolled under the Medicare program under title XVIII or under a
State Medicaid plan under title XIX or under a waiver of such plan.

“(3) WHOLESALE ACQUISITION COST.—The term ‘wholesale acquisition cost’ has the meaning given that term in section 1847A(c)(6)(B).

“(b) REPORT.—

“(1) REPORT REQUIRED.—The manufacturer of a qualifying drug shall submit a report to the Secretary if, with respect to the qualifying drug—

“(A) there is an increase in the price of the qualifying drug that results in an increase in the wholesale acquisition cost of that drug that is equal to—

“(i) 10 percent or more within a 12-month period beginning on or after January 1, 2019; or

“(ii) 25 percent or more within a 36-month period beginning on or after January 1, 2019;

“(B) the estimated price of the qualifying drug or spending per individual or per user of such drug (as estimated by the Secretary) for the applicable year (or per course of treatment in such applicable year as determined by the Sec-
(C) there was an increase in the price of the qualifying drug that resulted in an increase in the wholesale acquisition cost of that drug that is equal to—

“(i) 10 percent or more within a 12-month period that begins and ends during the 5-year period preceding January 1, 2021; or

“(ii) 25 percent or more within a 36-month period that begins and ends during the 5-year period preceding January 1, 2021.

“(2) REPORT DEADLINE.—Each report described in paragraph (1) shall be submitted to the Secretary—

“(A) in the case of a report with respect to an increase in the price of a qualifying drug that occurs during the period beginning on January 1, 2019, and ending on the day that is 60 days after the date of the enactment of this section, not later than 90 days after such date of enactment;
“(B) in the case of a report with respect to an increase in the price of a qualifying drug that occurs after the period described in subparagraph (A), not later than 30 days prior to the planned effective date of such price increase for such qualifying drug;

“(C) in the case of a report with respect to a qualifying drug that meets the criteria under paragraph (1)(B), not later than 30 days after such drug meets such criteria; and

“(D) in the case of a report with respect to an increase in the price of a qualifying drug that occurs during a 12-month or 36-month period described in paragraph (1)(C), not later than April 1, 2021.

“(c) CONTENTS.—A report under subsection (b), consistent with the standard for disclosures described in section 213.3(d) of title 12, Code of Federal Regulations (as in effect on the date of enactment of this section), shall, at a minimum, include—

“(1) with respect to the qualifying drug—

“(A) the percentage by which the manufacturer will raise the wholesale acquisition cost of the drug within the 12-month period or 36-month period as described in subsection
(b)(1)(A)(i), (b)(1)(A)(ii), (b)(1)(C)(i), or (b)(1)(C)(ii), as applicable, and the effective date of such price increase or the cost associated with a qualifying drug if such drug meets the criteria under subsection (b)(1)(B) and the effective date at which such drug meets such criteria;

“(B) an explanation for, and description of, each price increase for such drug that will occur during the 12-month period or the 36-month period described in subsection (b)(1)(A)(i), (b)(1)(A)(ii), (b)(1)(C)(i), or (b)(1)(C)(ii), as applicable;

“(C) an explanation for, and description of, the cost associated with a qualifying drug if such drug meets the criteria under subsection (b)(1)(B), as applicable;

“(D) if known and different from the manufacturer of the qualifying drug, the identity of—

“(i) the sponsor or sponsors of any investigational new drug applications under section 505(i) of the Federal Food, Drug, and Cosmetic Act for clinical investigations with respect to such drug, for which the full reports are submitted as part of the application—
“(I) for approval of the drug under section 505 of such Act; or

“(II) for licensure of the drug under section 351 of the Public Health Service Act; and

“(ii) the sponsor of an application for the drug approved under such section 505 of the Federal Food, Drug, and Cosmetic Act or licensed under section 351 of the Public Health Service Act;

“(E) a description of the history of the manufacturer’s price increases for the drug since the approval of the application for the drug under section 505 of the Federal Food, Drug, and Cosmetic Act or the issuance of the license for the drug under section 351 of the Public Health Service Act, or since the manufacturer acquired such approved application or license, if applicable;

“(F) the current wholesale acquisition cost of the drug;

“(G) the total expenditures of the manufacturer on—

“(i) materials and manufacturing for such drug;
“(ii) acquiring patents and licensing for such drug; and

“(iii) purchasing or acquiring such drug from another manufacturer, if applicable;

“(H) the percentage of total expenditures of the manufacturer on research and development for such drug that was derived from Federal funds;

“(I) the total expenditures of the manufacturer on research and development for such drug that is necessary to demonstrate that it meets applicable statutory standards for approval under section 505 of the Federal Food, Drug, and Cosmetic Act or licensure under section 351 of the Public Health Service Act, as applicable;

“(J) the total expenditures of the manufacturer on pursuing new or expanded indications or dosage changes for such drug under section 505 of the Federal Food, Drug, and Cosmetic Act or section 351 of the Public Health Service Act;

“(K) the total expenditures of the manufacturer on carrying out postmarket requirements related to such drug, including under section
505(o)(3) of the Federal Food, Drug, and Cosmetic Act;

“(L) the total revenue and the net profit generated from the qualifying drug for each calendar year since the approval of the application for the drug under section 505 of the Federal Food, Drug, and Cosmetic Act or the issuance of the license for the drug under section 351 of the Public Health Service Act, or since the manufacturer acquired such approved application or license; and

“(M) the total costs associated with marketing and advertising for the qualifying drug;

“(2) with respect to the manufacturer—

“(A) the total revenue and the net profit of the manufacturer for each of the 12-month period described in subsection (b)(1)(A)(i) or (b)(1)(C)(i) or the 36-month period described in subsection (b)(1)(A)(ii) or (b)(1)(C)(ii), as applicable;

“(B) all stock-based performance metrics used by the manufacturer to determine executive compensation for each of the 12-month periods described in subsection (b)(1)(A)(i) or (b)(1)(C)(i) or the 36-month periods described in
subsection (b)(1)(A)(ii) or (b)(1)(C)(ii), as applicable; and

“(C) any additional information the manufacturer chooses to provide related to drug pricing decisions, such as total expenditures on—

“(i) drug research and development; or

“(ii) clinical trials, including on drugs that failed to receive approval by the Food and Drug Administration; and

“(3) such other related information as the Secretary considers appropriate and as specified by the Secretary.

“(d) INFORMATION PROVIDED.—The manufacturer of a qualifying drug that is required to submit a report under subsection (b), shall ensure that such report and any explanation for, and description of, each price increase described in subsection (c)(1) shall be truthful, not misleading, and accurate.

“(e) CIVIL MONETARY PENALTY.—Any manufacturer of a qualifying drug that fails to submit a report for the drug as required by this section, following notification by the Secretary to the manufacturer that the manufacturer is not in compliance with this section, shall be subject to a civil monetary penalty of $75,000 for each day on which the violation continues.
“(f) False Information.—Any manufacturer that submits a report for a drug as required by this section that knowingly provides false information in such report is subject to a civil monetary penalty in an amount not to exceed $100,000 for each item of false information.

“(g) Public Posting.—

“(1) In General.—Subject to paragraph (4), the Secretary shall post each report submitted under subsection (b) on the public website of the Department of Health and Human Services the day the price increase of a qualifying drug is scheduled to go into effect.

“(2) Format.—In developing the format in which reports will be publicly posted under paragraph (1), the Secretary shall consult with stakeholders, including beneficiary groups, and shall seek feedback from consumer advocates and readability experts on the format and presentation of the content of such reports to ensure that such reports are—

“(A) user-friendly to the public; and

“(B) written in plain language that consumers can readily understand.

“(3) List.—In addition to the reports submitted under subsection (b), the Secretary shall also post a list of each qualifying drug with respect to which the
manufacturer was required to submit such a report in the preceding year and whether such manufacturer was required to submit such report based on a qualifying price increase or whether such drug meets the criteria under subsection (b)(1)(B).

“(4) PROTECTED INFORMATION.—In carrying out this section, the Secretary shall enforce applicable law concerning the protection of confidential commercial information and trade secrets.

“SEC. 1150D. ANNUAL REPORT TO CONGRESS.

“(a) IN GENERAL.—Subject to subsection (b), the Secretary shall submit to the Committees on Energy and Commerce and Ways and Means of the House of Representatives and the Committees on Health, Education, Labor, and Pensions and Finance of the Senate, and post on the public website of the Department of Health and Human Services in a way that is user-friendly to the public and written in plain language that consumers can readily understand, an annual report—

“(1) summarizing the information reported pursuant to section 1150C;

“(2) including copies of the reports and supporting detailed economic analyses submitted pursuant to such section;
“(3) detailing the costs and expenditures incurred by the Department of Health and Human Services in carrying out section 1150C; and

“(4) explaining how the Department of Health and Human Services is improving consumer and provider information about drug value and drug price transparency.

“(b) PROTECTED INFORMATION.—In carrying out this section, the Secretary shall enforce applicable law concerning the protection of confidential commercial information and trade secrets.”.