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(Original Signature of Member)

116TH CONGRESS
1ST SESSION

H. R.

To amend titles XI and XVIII of the Social Security Act to provide for drug manufacturer price transparency, to require certain manufacturers to report on product samples provided to certain health care providers, and for other purposes.

IN THE HOUSE OF REPRESENTATIVES

Mr. NEAL (for himself and [see ATTACHED LIST of cosponsors]) introduced the following bill; which was referred to the Committee on

A BILL

To amend titles XI and XVIII of the Social Security Act to provide for drug manufacturer price transparency, to require certain manufacturers to report on product samples provided to certain health care providers, and for other purposes.

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

1 **SECTION 1. SHORT TITLE.**

2 This Act may be cited as the “Prescription Drug
3 Sunshine, Transparency, Accountability and Reporting
4 Act” or the “Prescription Drug STAR Act”.

5 **SEC. 2. DRUG MANUFACTURER PRICE TRANSPARENCY.**

6 (a) IN GENERAL.—Title XI of the Social Security Act
7 (42 U.S.C. 1301 et seq.) is amended by inserting after
8 section 1128K the following new section:

9 **“SEC. 1128L. DRUG MANUFACTURER PRICE TRANS-**
10 **PARENCY.**

11 “(a) IN GENERAL.—With respect to each year, begin-
12 ning with 2021, the Secretary shall, at least once during
13 such year, determine if there is a triggered SPIKE in-
14 crease (in accordance with subsection (b)) with respect to
15 an applicable drug (as defined in subsection (f)(1)). If the
16 Secretary determines, with respect to a year, there is such
17 an increase with respect to an applicable drug, the manu-
18 facturer of the applicable drug shall submit to the Sec-
19 retary the justification described in subsection (c), subject
20 to subsection (b)(4), for each such triggered SPIKE in-
21 crease in accordance with the timing described in sub-
22 section (d)).

23 “(b) TRIGGERED SPIKE INCREASE.—

24 “(1) IN GENERAL.—A triggered SPIKE in-
25 crease occurs, with respect an applicable drug and
26 year (beginning with 2021 and referred to in this

1 paragraph as the ‘applicable year’), in any of the fol-
2 lowing cases:

3 “(A) If there is at least a 10 percent (or
4 \$10,000) cumulative increase with respect to
5 the wholesale acquisition cost (or alternative
6 cost measure specified by the Secretary under
7 paragraph (3)) of such drug during a calendar-
8 year period beginning and ending within the
9 lookback period that is the 5-year period pre-
10 ceeding such applicable year.

11 “(B) If there is at least a 25 percent (or
12 \$25,000) cumulative increase with respect to
13 the wholesale acquisition cost (or such alter-
14 native cost measure) of such drug during any
15 three-calendar-year period beginning and end-
16 ing within such lookback period.

17 “(C) In the case of such a drug that is
18 first covered under title XVIII with respect to
19 such applicable year, if the estimated cost or
20 spending under such title per individual or per
21 user of such drug (as estimated by the Sec-
22 retary) for such applicable year (or per course
23 of treatment in such applicable year, as defined
24 by the Secretary) is at least \$26,000.

1 “(2) INDEXING DOLLAR AMOUNTS.—The dollar
2 amounts applied under paragraph (1) for 2022 and
3 each subsequent year shall be the dollar amounts
4 specified in such paragraph for the previous year in-
5 creased by the annual percentage increase in the
6 consumer price index (all items; U.S. city average)
7 as of September of such previous year. If any
8 amount established under paragraph (1), after appli-
9 cation of this paragraph, for a year is not a multiple
10 of \$10, it shall be rounded to the nearest multiple
11 of \$10.

12 “(3) ALTERNATIVE TO WAC.—The Secretary
13 may, for purposes of making determinations under
14 paragraph (1), in addition to using the wholesale ac-
15 quisition cost for an applicable drug, use alternative
16 cost measures of such drug, or use such alternative
17 cost measure if the wholesale acquisition cost is not
18 available.

19 “(4) EXCEPTION.—A justification under sub-
20 section (c) shall not be required for a triggered
21 SPIKE increase described in paragraph (1) of an
22 applicable drug of a manufacturer if—

23 “(A) there is any portion of the lookback
24 period described in the respective subparagraph
25 of such paragraph for such increase that is in-

1 cluded within the lookback period for another
2 triggered SPIKE increase (or combination of
3 such increases) for which a justification is made
4 under this section for such drug by such manu-
5 facturer; or

6 “(B) such increase is less than the whole-
7 sale acquisition cost (or alternative cost meas-
8 ure specified by the Secretary under paragraph
9 (3)) of such drug during the calendar-year pe-
10 riod described in paragraph (1)(A) or the three-
11 calendar-year period described in paragraph
12 (1)(B), as applicable, for such increase, in-
13 creased by the percentage increase in the con-
14 sumer price index for all urban consumers (all
15 items; United States city average) for the 12-
16 month period ending six months prior to the
17 calendar-year period so described and for the
18 36-month period ending six months prior to the
19 three-calendar-year period so described, respec-
20 tively.

21 “(5) UNIT DETERMINATION.—For purposes of
22 determining the wholesale acquisition cost in car-
23 rying out this section, the Secretary shall determine
24 a unit (such as a unit size) to apply.

1 “(6) PUBLIC POSTING.—Beginning with respect
2 to 2021, the Secretary shall publicly post on the
3 Internet website of the Department of Health and
4 Human Services—

5 “(A) alternative percentages, dollar
6 amounts, and lookback periods that, if applied
7 under paragraph (1), would be projected to in-
8 crease the number of applicable drugs for which
9 a triggered SPIKE increase would occur for
10 such year; and

11 “(B) the number of applicable drugs for
12 which a triggered SPIKE increase would occur
13 for such year of such an alternative percentage,
14 dollar amount, or period were applied for such
15 year.

16 “(c) JUSTIFICATION DESCRIBED.—

17 “(1) IN GENERAL.—The justification described
18 in this subsection, with respect to a triggered
19 SPIKE increase described in subsection (b)(1) of an
20 applicable drug of a manufacturer, is—

21 “(A) all of the information described in
22 paragraph (2);

23 “(B) all of the information and supporting
24 documentation described in paragraph (3), as
25 applicable to the increase and drug; and

1 “(C) a certification described in paragraph
2 (4).

3 “(2) REQUIRED INFORMATION.—For purposes
4 of paragraph (1), the information described in this
5 paragraph is the following:

6 “(A) The individual factors that have con-
7 tributed to the increase in the wholesale acqui-
8 sition cost.

9 “(B) An explanation of the role of each
10 factor in contributing to such increase.

11 “(3) INFORMATION AS APPLICABLE.—For pur-
12 poses of paragraph (1), the information and sup-
13 porting documentation described in this paragraph is
14 the following, as applicable to the increase of the
15 drug:

16 “(A) Total expenditures of the manufac-
17 turer on—

18 “(i) materials and manufacturing for
19 such drug;

20 “(ii) acquiring patents and licensing
21 for each drug of the manufacturer; and

22 “(iii) costs to purchase or acquire the
23 drug from another company, if applicable.

24 “(B) The percentage of total expenditures
25 of the manufacturer on research and develop-

1 ment for such drug that was derived from Fed-
2 eral funds.

3 “(C) The total expenditures of the manu-
4 facturer on research and development for such
5 drug.

6 “(D) The total revenue and net profit gen-
7 erated from the applicable drug for each cal-
8 endar year since drug approval.

9 “(E) The total costs associated with mar-
10 keting and advertising for the applicable drug.

11 “(F) Additional information specific to the
12 manufacturer of the applicable drug, such as—

13 “(i) the total revenue and net profit of
14 the manufacturer for the period of such in-
15 crease, as determined by the Secretary;

16 “(ii) metrics used to determine execu-
17 tive compensation;

18 “(iii) total expenditures on—

19 “(I) drug research and develop-
20 ment; or

21 “(II) clinical trials on drugs that
22 failed to receive approval by the Food
23 and Drug Administration; and

1 “(iv) any additional information re-
2 lated to drug pricing decisions of the man-
3 ufacturer.

4 “(G) Any other relevant information and
5 supporting documentation necessary to justify
6 the triggering SPIKE increase.

7 “(H) Any other relevant information and
8 supporting documentation, as specified by the
9 Secretary.

10 “(4) CERTIFICATION.—For purposes of para-
11 graph (1), the certification described in this para-
12 graph is a certification, that all such information
13 and documentation is accurate and complete, by one
14 of the following:

15 “(A) The chief executive officer of the
16 manufacturer.

17 “(B) The chief financial officer of the
18 manufacturer.

19 “(C) An individual who has delegated au-
20 thority to sign for, and who reports directly to,
21 such chief executive officer or chief financial of-
22 ficer.

23 “(d) TIMING.—

24 “(1) NOTIFICATION.—Not later than 60 days
25 after the date on which the Secretary makes the de-

1 termination that there is a triggering SPIKE in-
2 crease with respect to an applicable drug, the Sec-
3 retary shall notify the manufacturer of the applica-
4 ble drug of such determination.

5 “(2) SUBMISSION OF JUSTIFICATION.—Not
6 later than 90 days after the date on which a manu-
7 facturer receives a notification under paragraph (1),
8 subject to subsection (b)(4), the manufacturer shall
9 submit to the Secretary the justification required
10 under subsection (a), including a summary of such
11 justification, in a form and manner specified by the
12 Secretary. In specifying such form, with respect to
13 the summary required under the previous sentence,
14 the Secretary shall provide that such summary shall
15 be in an easily understandable format, as specified
16 by the Secretary, and shall permit the manufacturer
17 to exclude proprietary information from such sum-
18 mary.

19 “(3) POSTING ON INTERNET WEBSITE.—Not
20 later than 30 days after receiving the complete jus-
21 tification under paragraph (2), the Secretary shall
22 post on the Internet website of the Centers for Medi-
23 care & Medicaid Services the summary included for
24 such justification.

25 “(e) PENALTIES.—

1 “(1) FAILURE TO SUBMIT TIMELY JUSTIFICA-
2 TION.—If the Secretary determines that a manufac-
3 turer has failed to submit a justification as required
4 under this section, including in accordance with the
5 timing and form required, with respect to an appli-
6 cable drug, the Secretary shall apply a civil mone-
7 etary penalty in an amount of \$10,000 for each day
8 the manufacturer has failed to submit such justifica-
9 tion as so required.

10 “(2) FALSE INFORMATION.—Any manufacturer
11 that submits a justification under this section that
12 knowingly provides false information in such jus-
13 tification is subject to a civil monetary penalty in an
14 amount not to exceed \$100,000 for each item of
15 false information.

16 “(3) APPLICATION OF PROCEDURES.—The pro-
17 visions of section 1128A (other than subsections (a)
18 and (b)) shall apply to a civil monetary penalty
19 under this subsection in the same manner as such
20 provisions apply to a penalty or proceeding under
21 section 1128A(a). Civil monetary penalties imposed
22 under this subsection are in addition to other pen-
23 alties as may be prescribed by law.

24 “(f) DEFINITIONS.—In this section:

25 “(1) APPLICABLE DRUG.—

1 “(A) IN GENERAL.—Subject to subpara-
2 graph (B), the term ‘applicable drug’ means,
3 with respect to a lookback period described in
4 subsection (b)(1), a covered outpatient drug (as
5 defined in paragraph (2) of section 1927(k),
6 without application of paragraph (3) of such
7 section) that is covered under title XVIII and
8 is not a low cost drug.

9 “(B) EXCLUSION OF LOW COST DRUGS.—
10 For purposes of subparagraph (A), not later
11 than January 1, 2021, the Secretary shall
12 specify a threshold (such as a cost or spending
13 threshold) for identifying (and shall identify)
14 low cost drugs to be excluded from the defini-
15 tion of the term ‘applicable drug’, such as a
16 drug that has a wholesale acquisition cost of
17 less than \$10 per unit or less than \$100 in av-
18 erage estimated expenditures under title XVIII
19 per individual per year or per user of such drug
20 per year. For purposes of this section, a drug
21 shall not be considered specified as a low cost
22 drug for a lookback period described in sub-
23 section (b)(1) with respect to a year unless such
24 drug is identified as being below the specified

1 threshold for the entirety of the lookback pe-
2 riod.

3 “(2) MANUFACTURER.—The term ‘manufac-
4 turer’ has the meaning given that term in section
5 1847A(c)(6)(A).

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

9 (b) REPORTING TO THE SECRETARY OF THE TREAS-
10 URY.—

11 (1) IN GENERAL.—Subpart A of part III of
12 subchapter A of chapter 61 of the Internal Revenue
13 Code of 1986 is amended by inserting after section
14 6039J the following new section:

15 **“SEC. 6039K. DRUG PRICE SPIKE INCREASE REPORTING.**

16 “Each manufacturer (within the meaning of section
17 1128L of the Social Security Act) shall file a return (as
18 such time and in such form and manner as the Secretary
19 may provide) showing for each year with respect to which
20 such section applies all information and supporting docu-
21 mentation and the certification included within a justifica-
22 tion reported by the manufacturer under subsection (c)(1)
23 of such section.”.

24 (2) CLERICAL AMENDMENT.—The table of sec-
25 tions for subpart A of part III of subchapter A of

1 chapter 61 of such Code is amended by inserting
2 after the item relating to section 6039J the fol-
3 lowing new item:

“Sec. 6039K. Drug price SPIKE increase reporting.”.

4 **SEC. 3. REQUIREMENT FOR MANUFACTURERS OF CERTAIN**
5 **DRUGS, DEVICES, BIOLOGICALS, AND MED-**
6 **ICAL SUPPLIES TO REPORT ON PRODUCT**
7 **SAMPLES PROVIDED TO CERTAIN HEALTH**
8 **CARE PROVIDERS.**

9 (a) IN GENERAL.—Section 1128G(a) of the Social
10 Security Act (42 U.S.C. 1320a–7h(a)) is amended by add-
11 ing at the end the following new paragraph:

12 “(3) CERTAIN PRODUCT SAMPLES.—

13 “(A) IN GENERAL.—In addition to the re-
14 quirements under paragraphs (1)(A) and (2),
15 on the 90th day of each calendar year (begin-
16 ning with 2023), any applicable manufacturer
17 that provides a payment or other transfer of
18 value that is a product sample described in sub-
19 paragraph (B) to any covered recipient (or to
20 an entity or individual at the request of, or des-
21 ignated on behalf of, such a covered recipient)
22 shall submit to the Secretary, in such electronic
23 form as the Secretary shall require, the fol-
24 lowing information (aggregated per each drug,

1 device, biological, or medical supply, as applica-
2 ble) with respect to the preceding calendar year:

3 “(i) The total quantity of all such
4 payments or other transfers of value pro-
5 vided to all covered recipients.

6 “(ii) The total value of all such pay-
7 ments or other transfers of value provided
8 to all covered recipients.

9 “(iii) If applicable, information de-
10 scribed in clauses (vii) and (viii) of para-
11 graph (1)(A) with respect to such a pay-
12 ment or other transfer of value.

13 “(B) PRODUCT SAMPLE DESCRIBED.—For
14 purposes of subparagraph (A), a product sam-
15 ple described in this subparagraph is a product
16 sample that is not intended to be sold and is in-
17 tended for patient use.”.

18 (b) PUBLIC AVAILABILITY OF INFORMATION.—Sec-
19 tion 1128G(e)(1)(C)(ii) of the Social Security Act (42
20 U.S.C. 1320a–7h(e)(1)(C)(ii)) is amended—

21 (1) by striking “(ii) contains” and inserting
22 “(ii)(I) with respect to information that is not infor-
23 mation submitted under paragraph (3) of subsection
24 (a), contains”;

1 (2) by striking “, as applicable;” and inserting
2 “, as applicable; and”;

3 (3) by adding at the end the following new sub-
4 clause:

5 “(II) with respect to information sub-
6 mitted under paragraph (3) of subsection
7 (a), contains information that is presented
8 by the name of the applicable manufac-
9 turer, the total amount of all payments or
10 other transfers of value described in such
11 paragraph provided to all covered recipi-
12 ents, the total value of all such payments
13 or other transfers of value provided to all
14 covered recipients, and the name of the
15 covered drug, device, biological, or medical
16 supply, as applicable;”.

17 (c) CONFORMING AMENDMENT.—Section
18 1128G(e)(10)(B)(ii) of the Social Security Act (42 U.S.C.
19 1320a–7h(e)(10)(B)(ii)) is amended by striking “Product
20 samples” and inserting “Except for purposes of paragraph
21 (3) of subsection (a), product samples”.

22 (d) REPORTING TO THE SECRETARY OF THE TREAS-
23 URY.—

24 (1) IN GENERAL.—Subpart A of part III of
25 subchapter A of chapter 61 of the Internal Revenue

1 Code of 1986, as amended by section 2, is further
2 amended by inserting after section 6039K the fol-
3 lowing new section:

4 **“SEC. 6039L. PRODUCT SAMPLES OF APPLICABLE MANU-
5 FACTURERS.**

6 “Each applicable manufacturer (within the meaning
7 of section 1128G(a)(3) of the Social Security Act) shall
8 file a return (as such time and in such form and manner
9 as the Secretary may provide) showing for each year to
10 which such section applies—

11 “(1) the amount described in section
12 1128G(a)(3)(A)(ii) of such Act with respect to such
13 year, and

14 “(2) the portion of such amount for which a de-
15 duction was claimed under section 162.”.

16 (2) CLERICAL AMENDMENT.—The table of sec-
17 tions for subpart A of part III of subchapter A of
18 chapter 61 of such Code, as amended by section 2,
19 is further amended by inserting after the item relat-
20 ing to section 6039K the following new item:

“Sec. 6039L. Product samples of applicable manufacturers.”.

21 **SEC. 4. ANALYSIS AND REPORT ON INPATIENT HOSPITAL
22 DRUG COSTS.**

23 (a) ANALYSIS.—The Secretary of Health and Human
24 Services shall conduct an analysis that, to the extent prac-
25 ticable—

1 (1) focuses on drugs that are furnished in the
2 inpatient setting;

3 (2) includes data on inpatient hospital drug
4 costs, Medicare spending, volume, and spending per
5 admission;

6 (3) considers trends in inpatient hospital drug
7 costs, such as trends by hospital size, classification
8 of urban or rural, whether the hospital is a teaching
9 hospital, or other categorization; and

10 (4) examines the impact of drug shortages on
11 services that are furnished in an inpatient hospital
12 setting.

13 In conducting such analysis, the Secretary may conduct
14 hospital surveys, use data from hospital cost reports, or
15 use other data as determined by the Secretary.

16 (b) REPORT.—Not later than January 1, 2021, the
17 Secretary shall submit to the Committee on Ways and
18 Means of the House of Representatives and the Finance
19 Committee of the Senate a report on drug costs in the
20 inpatient hospital setting, including the analyses described
21 in paragraphs (1) through (4) of subsection (a).

22 (c) FUNDING.—For purposes of carrying out this sec-
23 tion, there shall be transferred to the Secretary
24 \$3,000,000 from the Federal Hospital Insurance Trust

1 Fund under section 1817 of the Social Security Act (42
2 U.S.C. 1395i).

3 **SEC. 5. PUBLIC DISCLOSURE OF DRUG DISCOUNTS.**

4 Section 1150A of the Social Security Act (42 U.S.C.
5 1320b–23) is amended—

6 (1) in subsection (e), in the matter preceding
7 paragraph (1), by inserting “(other than as per-
8 mitted under subsection (e))” after “disclosed by the
9 Secretary”; and

10 (2) by adding at the end the following new sub-
11 section:

12 “(e) PUBLIC AVAILABILITY OF CERTAIN INFORMA-
13 TION.—

14 “(1) IN GENERAL.—In order to allow the com-
15 parison of PBMs’ ability to negotiate rebates, dis-
16 counts, and price concessions and the amount of
17 such rebates, discounts, and price concessions that
18 are passed through to plan sponsors, beginning Jan-
19 uary 1, 2020, the Secretary shall make available on
20 the Internet website of the Department of Health
21 and Human Services the information with respect to
22 the second preceding calendar year provided to the
23 Secretary on generic dispensing rates (as described
24 in paragraph (1) of subsection (b) and information
25 provided to the Secretary under paragraphs (2) and

1 (3) of such subsection that, as determined by the
2 Secretary, is with respect to each PBM.

3 “(2) AVAILABILITY OF DATA.—In carrying out
4 paragraph (1), the Secretary shall ensure the fol-
5 lowing:

6 “(A) CONFIDENTIALITY.—The information
7 described in such paragraph is displayed in a
8 manner that prevents the disclosure of informa-
9 tion on rebates, discounts, and price conces-
10 sions, with respect to an individual drug or an
11 individual plan.

12 “(B) CLASS OF DRUG.—The information
13 described in such paragraph is made available
14 by class of drug, using an existing classification
15 system, but only if the class contains such num-
16 ber of drugs, as specified by the Secretary, to
17 ensure confidentiality of proprietary informa-
18 tion or other information that is prevented to
19 be disclosed under subparagraph (A).”.

20 **SEC. 6. REQUIRING CERTAIN MANUFACTURERS TO REPORT**
21 **DRUG PRICING INFORMATION WITH RE-**
22 **SPECT TO DRUGS UNDER THE MEDICARE**
23 **PROGRAM.**

24 (a) IN GENERAL.—Section 1847A of the Social Secu-
25 rity Act (42 U.S.C. 1395w–3a) is amended—

1 (1) in subsection (b)—

2 (A) in paragraph (2)(A), by inserting “or
3 subsection (f)(2), as applicable” before the pe-
4 riod at the end;

5 (B) in paragraph (3), in the matter pre-
6 ceding subparagraph (A), by inserting “or sub-
7 section (f)(2), as applicable,” before “deter-
8 mined by”; and

9 (C) in paragraph (6)(A), in the matter
10 preceding clause (i), by inserting “or subsection
11 (f)(2), as applicable,” before “determined by”;
12 and

13 (2) in subsection (f)—

14 (A) by striking “For requirements” and
15 inserting the following:

16 “(1) IN GENERAL.—For requirements”; and

17 (B) by adding at the end the following new
18 paragraph:

19 “(2) MANUFACTURERS WITHOUT A REBATE
20 AGREEMENT UNDER TITLE XIX.—

21 “(A) IN GENERAL.—In the case of a man-
22 ufacturer of a drug or biological described in
23 subparagraph (C), (E), or (G) of section
24 1842(o)(1) or in clause (ii) or (iii) of section
25 1881(b)(14)(B) that does not have a rebate

1 agreement in effect under section 1927, for cal-
2 endar quarters beginning on or after January
3 1, 2020, such manufacturer shall report to the
4 Secretary the information described in sub-
5 section (b)(3)(A)(iii) of such section 1927 with
6 respect to such drug or biological in a time and
7 manner specified by the Secretary.

8 “(B) AUDIT.—Information reported under
9 subparagraph (A) is subject to audit by the In-
10 spector General of the Department of Health
11 and Human Services.

12 “(C) VERIFICATION.—The Secretary may
13 survey wholesalers and manufacturers that di-
14 rectly distribute drugs described in subpara-
15 graph (A), when necessary, to verify manufac-
16 turer prices and manufacturer’s average sales
17 prices (including wholesale acquisition cost) if
18 required to make payment reported under sub-
19 paragraph (A). The Secretary may impose a
20 civil monetary penalty in an amount not to ex-
21 ceed \$100,000 on a wholesaler, manufacturer,
22 or direct seller, if the wholesaler, manufacturer,
23 or direct seller of such a drug refuses a request
24 for information about charges or prices by the
25 Secretary in connection with a survey under

1 this subparagraph or knowingly provides false
2 information. The provisions of section 1128A
3 (other than subsections (a) (with respect to
4 amounts of penalties or additional assessments)
5 and (b)) shall apply to a civil money penalty
6 under this subparagraph in the same manner as
7 such provisions apply to a penalty or proceeding
8 under section 1128A(a).

9 “(D) CONFIDENTIALITY.—Notwith-
10 standing any other provision of law, information
11 disclosed by manufacturers or wholesalers
12 under this paragraph (other than the wholesale
13 acquisition cost for purposes of carrying out
14 this section) is confidential and shall not be dis-
15 closed by the Secretary in a form which dis-
16 closes the identity of a specific manufacturer or
17 wholesaler or prices charged for drugs by such
18 manufacturer or wholesaler, except—

19 “(i) as the Secretary determines to be
20 necessary to carry out this section (includ-
21 ing the determination and implementation
22 of the payment amount), or to carry out
23 section 1847B;

1 “(ii) to permit the Comptroller Gen-
2 eral to review the information provided;
3 and

4 “(iii) to permit the Director of the
5 Congressional Budget Office to review the
6 information provided.”.

7 (b) ENFORCEMENT.—Section 1847A such Act (42
8 U.S.C. 1395w-3a) is further amended—

9 (1) in subsection (d)(4)—

10 (A) in subparagraph (A), by striking “IN
11 GENERAL” and inserting “MISREPRESENTA-
12 TION”;

13 (B) in subparagraph (B), by striking “sub-
14 paragraph (B)” and inserting “subparagraph
15 (A), (B), or (C)”;

16 (C) by redesignating subparagraph (B) as
17 subparagraph (D); and

18 (D) by inserting after subparagraph (A)
19 the following new subparagraphs:

20 “(B) FAILURE TO PROVIDE TIMELY INFOR-
21 MATION.—If the Secretary determines that a
22 manufacturer described in subsection (f)(2) has
23 failed to report on information described in sec-
24 tion 1927(b)(3)(A)(iii) with respect to a drug or
25 biological in accordance with such subsection,

1 the Secretary shall apply a civil money penalty
2 in an amount of \$10,000 for each day the man-
3 ufacturer has failed to report such information
4 and such amount shall be paid to the Treasury.

5 “(C) FALSE INFORMATION.—Any manu-
6 facturer required to submit information under
7 subsection (f)(2) that knowingly provides false
8 information is subject to a civil money penalty
9 in an amount not to exceed \$100,000 for each
10 item of false information. Such civil money pen-
11 alties are in addition to other penalties as may
12 be prescribed by law.”; and

13 (2) in subsection (c)(6)(A), by striking the pe-
14 riod at the end and inserting “, except that, for pur-
15 poses of subsection (f)(2), the Secretary may, if the
16 Secretary determines appropriate, exclude repack-
17 agers of a drug or biological from such term.”.

18 (c) REPORT.—Not later than January 1, 2021, the
19 Inspector General of the Department of Health and
20 Human Services shall assess and submit to Congress a
21 report on the accuracy of average sales price information
22 submitted by manufacturers under section 1847A of the
23 Social Security Act (42 U.S.C. 1395w–3a). Such report
24 shall include any recommendations on how to improve the
25 accuracy of such information.