

111TH CONGRESS
1ST SESSION

S. 1239

To amend section 340B of the Public Health Service Act to revise and expand the drug discount program under that section to improve the provision of discounts on drug purchases for certain safety net providers.

IN THE SENATE OF THE UNITED STATES

JUNE 11, 2009

Mr. BINGAMAN (for himself, Mr. THUNE, and Mrs. GILLIBRAND) introduced the following bill; which was read twice and referred to the Committee on Health, Education, Labor, and Pensions

A BILL

To amend section 340B of the Public Health Service Act to revise and expand the drug discount program under that section to improve the provision of discounts on drug purchases for certain safety net providers.

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

3 **SECTION 1. SHORT TITLE.**

4 This Act may be cited as the “340B Program Im-
5 provement and Integrity Act of 2009”.

1 **SEC. 2. EXPANDED PARTICIPATION IN SECTION 340B PRO-**
2 **GRAM.**

3 (a) EXPANSION OF COVERED ENTITIES RECEIVING
4 DISCOUNTED PRICES.—Section 340B(a)(4) of the Public
5 Health Service Act (42 U.S.C. 256b(a)(4)) is amended by
6 adding at the end the following:

7 “(M) A children’s hospital excluded from
8 the Medicare prospective payment system pur-
9 suant to section 1886(d)(1)(B)(iii) of the Social
10 Security Act which would meet the require-
11 ments of subparagraph (L), including the dis-
12 proportionate share adjustment percentage re-
13 quirement under clause (ii) of such subpara-
14 graph, if the hospital were a subsection (d) hos-
15 pital as defined by section 1886(d)(1)(B) of the
16 Social Security Act.

17 “(N) An entity that is a critical access hos-
18 pital (as determined under section 1820(e)(2)
19 of the Social Security Act), and that meets the
20 requirements of subparagraph (L)(i).

21 “(O) An entity that is a rural referral cen-
22 ter, as defined by section 1886(d)(5)(C)(i) of
23 the Social Security Act, or a sole community
24 hospital, as defined by section
25 1886(d)(5)(C)(iii) of such Act, and that both
26 meets the requirements of subparagraph (L)(i)

1 and has a disproportionate share adjustment
2 percentage equal to or greater than 8 percent.”.

3 (b) EXTENSION OF DISCOUNTS TO INPATIENT
4 DRUGS.—Section 340B of the Public Health Service Act
5 (42 U.S.C. 256b) is amended—

6 (1) in subsection (a), by striking “outpatient”
7 each place that such appears in paragraphs (2), (5),
8 (7), and (9); and

9 (2) in subsection (b)—

10 (A) by striking “In this section” and in-
11 serting the following:

12 “(A) IN GENERAL.—In this section”; and

13 (B) by adding at the end the following:

14 “(B) COVERED DRUG.—In this section, the
15 term ‘covered drug’—

16 “(i) means a covered outpatient drug
17 (as defined in section 1927(k)(2) of the
18 Social Security Act); and

19 “(ii) includes, notwithstanding para-
20 graph (3)(A) of such section 1927(k), a
21 drug used in connection with an inpatient
22 or outpatient service provided by a hospital
23 described in subparagraph (L), (M), (N),
24 or (O) of subsection (a)(4) that is enrolled

1 to participate in the drug discount pro-
2 gram under this section.

3 “(C) PURCHASING ARRANGEMENTS FOR
4 INPATIENT DRUGS.—The Secretary shall ensure
5 that a hospital described in subparagraph (L),
6 (M), (N), or (O) of subsection (a)(4) that is en-
7 rolled to participate in the drug discount pro-
8 gram under this section shall have multiple op-
9 tions for purchasing covered drugs for inpa-
10 tients including by utilizing a group purchasing
11 organization or other group purchasing ar-
12 rangement, establishing and utilizing its own
13 group purchasing program, purchasing directly
14 from a manufacturer, and any other purchasing
15 arrangements that the Secretary may deem ap-
16 propriate to ensure access to drug discount
17 pricing under this section for inpatient drugs
18 taking into account the particular needs of
19 small and rural hospitals.”.

20 (c) PROHIBITION ON GROUP PURCHASING ARRANGE-
21 MENTS.—Section 340B(a) of the Public Health Service
22 Act (42 U.S.C. 256b(a)) is amended—

23 (1) in paragraph (4)(L)—

24 (A) in clause (i), by adding “and” at the
25 end;

1 (B) in clause (ii), by striking “; and” and
2 inserting a period; and

3 (C) by striking clause (iii); and
4 (2) in paragraph (5)—

5 (A) by redesignating subparagraphs (C)
6 and (D) as subparagraphs (D) and (E); respec-
7 tively; and

8 (B) by inserting after subparagraph (B),
9 the following:

10 “(C) PROHIBITING THE USE OF GROUP
11 PURCHASING ARRANGEMENTS.—

12 “(i) IN GENERAL.—A hospital de-
13 scribed in subparagraphs (L), (M), (N), or
14 (O) of paragraph (4) shall not obtain cov-
15 ered outpatient drugs through a group
16 purchasing organization or other group
17 purchasing arrangement, except as per-
18 mitted or provided for pursuant to clauses
19 (ii) or (iii).

20 “(ii) INPATIENT DRUGS.—Clause (i)
21 shall not apply to drugs purchased for in-
22 patient use.

23 “(iii) EXCEPTIONS.—The Secretary
24 shall establish reasonable exceptions to
25 clause (i)—

1 “(I) with respect to a covered
2 outpatient drug that is unavailable to
3 be purchased through the program
4 under this section due to a drug
5 shortage problem, manufacturer non-
6 compliance, or any other circumstance
7 beyond the hospital’s control;

8 “(II) to facilitate generic substi-
9 tution when a generic covered out-
10 patient drug is available at a lower
11 price; or

12 “(III) to reduce in other ways
13 the administrative burdens of man-
14 aging both inventories of drugs sub-
15 ject to this section and inventories of
16 drugs that are not subject to this sec-
17 tion, so long as the exceptions do not
18 create a duplicate discount problem in
19 violation of subparagraph (A) or a di-
20 version problem in violation of sub-
21 paragraph (B).”.

22 (d) MEDICAID CREDITS ON INPATIENT DRUGS.—
23 Section 340B(a)(5) of the Public Health Service Act (42
24 U.S.C. 256b(a)(5)) is amended by adding at the end the
25 following:

1 “(E) MEDICAID CREDITS.—Not later than
2 90 days after the date of filing of the hospital’s
3 most recently filed Medicare cost report, the
4 hospital shall issue a credit as determined by
5 the Secretary to the State Medicaid program
6 for inpatient covered drugs provided to Med-
7 icaid recipients.”.

8 (e) INTEGRITY IMPROVEMENTS.—Subsection (c) of
9 section 340B of the Public Health Service Act (42 U.S.C.
10 256b(c)) is amended to read as follows:

11 “(c) IMPROVEMENTS IN PROGRAM INTEGRITY.—

12 “(1) MANUFACTURER COMPLIANCE.—

13 “(A) IN GENERAL.—From amounts appro-
14 priated under paragraph (4), the Secretary
15 shall provide for improvements in compliance by
16 manufacturers with the requirements of this
17 section in order to prevent overcharges and
18 other violations of the discounted pricing re-
19 quirements specified in this section.

20 “(B) IMPROVEMENTS.—The improvements
21 described in subparagraph (A) shall include the
22 following:

23 “(i) The development of a system to
24 enable the Secretary to verify the accuracy
25 of ceiling prices calculated by manufactur-

1 ers under subsection (a)(1) and charged to
2 covered entities, which shall include the
3 following:

4 “(I) Developing and publishing
5 through an appropriate policy or regu-
6 latory issuance, precisely defined
7 standards and methodology for the
8 calculation of ceiling prices under
9 such subsection.

10 “(II) Comparing regularly the
11 ceiling prices calculated by the Sec-
12 retary with the quarterly pricing data
13 that is reported by manufacturers to
14 the Secretary.

15 “(III) Performing spot checks of
16 sales transactions by covered entities.

17 “(IV) Inquiring into the cause of
18 any pricing discrepancies that may be
19 identified and either taking, or requir-
20 ing manufacturers to take, such cor-
21 rective action as is appropriate in re-
22 sponse to such price discrepancies.

23 “(ii) The establishment of procedures
24 for manufacturers to issue refunds to cov-
25 ered entities in the event that there is an

1 overcharge by the manufacturers, including
2 the following:

3 “(I) Providing the Secretary with
4 an explanation of why and how the
5 overcharge occurred, how the refunds
6 will be calculated, and to whom the
7 refunds will be issued.

8 “(II) Oversight by the Secretary
9 to ensure that the refunds are issued
10 accurately and within a reasonable pe-
11 riod of time, both in routine instances
12 of retroactive adjustment to relevant
13 pricing data and exceptional cir-
14 cumstances such as erroneous or in-
15 tentional overcharging for covered
16 drugs.

17 “(iii) The provision of access through
18 the Internet website of the Department of
19 Health and Human Services to the applica-
20 ble ceiling prices for covered drugs as cal-
21 culated and verified by the Secretary in ac-
22 cordance with this section, in a manner
23 (such as through the use of password pro-
24 tection) that limits such access to covered
25 entities and adequately assures security

1 and protection of privileged pricing data
2 from unauthorized re-disclosure.

3 “(iv) The development of a mecha-
4 nism by which—

5 “(I) rebates and other discounts
6 provided by manufacturers to other
7 purchasers subsequent to the sale of
8 covered drugs to covered entities are
9 reported to the Secretary; and

10 “(II) appropriate credits and re-
11 funds are issued to covered entities if
12 such discounts or rebates have the ef-
13 fect of lowering the applicable ceiling
14 price for the relevant quarter for the
15 drugs involved.

16 “(v) Selective auditing of manufactur-
17 ers and wholesalers to ensure the integrity
18 of the drug discount program under this
19 section.

20 “(vi) The imposition of sanctions in
21 the form of civil monetary penalties,
22 which—

23 “(I) shall be assessed according
24 to standards established in regulations
25 to be promulgated by the Secretary

1 within 180 days of the date of enact-
2 ment of the 340B Program Improve-
3 ment and Integrity Act of 2009;

4 “(II) shall not exceed \$5,000 for
5 each instance of overcharging a cov-
6 ered entity that may have occurred;
7 and

8 “(III) shall apply to any manu-
9 facturer with an agreement under this
10 section that knowingly and inten-
11 tionally charges a covered entity a
12 price for purchase of a drug that ex-
13 ceeds the maximum applicable price
14 under subsection (a)(1).

15 “(2) COVERED ENTITY COMPLIANCE.—

16 “(A) IN GENERAL.—From amounts appro-
17 priated under paragraph (4), the Secretary
18 shall provide for improvements in compliance by
19 covered entities with the requirements of this
20 section in order to prevent diversion and viola-
21 tions of the duplicate discount provision and
22 other requirements specified under subsection
23 (a)(5).

1 “(B) IMPROVEMENTS.—The improvements
2 described in subparagraph (A) shall include the
3 following:

4 “(i) The development of procedures to
5 enable and require covered entities to regu-
6 larly update (at least annually) the infor-
7 mation on the Internet website of the De-
8 partment of Health and Human Services
9 relating to this section.

10 “(ii) The development of a system for
11 the Secretary to verify the accuracy of in-
12 formation regarding covered entities that is
13 listed on the website described in clause
14 (i).

15 “(iii) The development of more de-
16 tailed guidance describing methodologies
17 and options available to covered entities for
18 billing covered drugs to State Medicaid
19 agencies in a manner that avoids duplicate
20 discounts pursuant to subsection (a)(5)(A).

21 “(iv) The establishment of a single,
22 universal, and standardized identification
23 system by which each covered entity site
24 can be identified by manufacturers, dis-
25 tributors, covered entities, and the Sec-

1 retary for purposes of facilitating the or-
2 dering, purchasing, and delivery of covered
3 drugs under this section, including the
4 processing of chargebacks for such drugs.

5 “(v) The imposition of sanctions, in
6 appropriate cases as determined by the
7 Secretary, additional to those to which cov-
8 ered entities are subject under subpara-
9 graph (a)(5)(E), through one or more of
10 the following actions:

11 “(I) Where a covered entity
12 knowingly and intentionally violates
13 subparagraph (a)(5)(B), the covered
14 entity shall be required to pay a mon-
15 etary penalty to a manufacturer or
16 manufacturers in the form of interest
17 on sums for which the covered entity
18 is found liable under paragraph
19 (a)(5)(E), such interest to be com-
20 pounded monthly and equal to the
21 current short term interest rate as de-
22 termined by the Federal Reserve for
23 the time period for which the covered
24 entity is liable.

1 “(II) Where the Secretary deter-
2 mines a violation of subparagraph
3 (a)(5)(B) was systematic and egre-
4 gious as well as knowing and inten-
5 tional, removing the covered entity
6 from the drug discount program
7 under this section and disqualifying
8 the entity from re-entry into such pro-
9 gram for a reasonable period of time
10 to be determined by the Secretary.

11 “(III) Referring matters to ap-
12 propriate Federal authorities within
13 the Food and Drug Administration,
14 the Office of Inspector General of De-
15 partment of Health and Human Serv-
16 ices, or other Federal agencies for
17 consideration of appropriate action
18 under other Federal statutes, such as
19 the Prescription Drug Marketing Act.

20 “(3) ADMINISTRATIVE DISPUTE RESOLUTION
21 PROCESS.—

22 “(A) IN GENERAL.—Not later than 180
23 days after the date of enactment of the 340B
24 Program Improvement and Integrity Act of
25 2009, the Secretary shall promulgate regula-

1 tions to establish and implement an administra-
2 tive process for the resolution of claims by cov-
3 ered entities that they have been overcharged
4 for drugs purchased under this section, and
5 claims by manufacturers, after the conduct of
6 audits as authorized by subsection (a)(5)(D), of
7 violations of subsections (a)(5)(A) or (a)(5)(B),
8 including appropriate procedures for the provi-
9 sion of remedies and enforcement of determina-
10 tions made pursuant to such process through
11 mechanisms and sanctions described in para-
12 graphs (1)(B) and (2)(B).

13 “(B) DEADLINE AND PROCEDURES.—Reg-
14 ulations promulgated by the Secretary under
15 subparagraph (A) shall—

16 “(i) designate or establish a decision-
17 making official or decision-making body
18 within the Department of Health and
19 Human Services to be responsible for re-
20 viewing and finally resolving claims by cov-
21 ered entities that they have been charged
22 prices for covered drugs in excess of the
23 ceiling price described in subsection (a)(1),
24 and claims by manufacturers that viola-

1 tions of subsection (a)(5)(A) or (a)(5)(B)
2 have occurred;

3 “(ii) establish such deadlines and pro-
4 cedures as may be necessary to ensure that
5 claims shall be resolved fairly, efficiently,
6 and expeditiously;

7 “(iii) establish procedures by which a
8 covered entity may discover and obtain
9 such information and documents from
10 manufacturers and third parties as may be
11 relevant to demonstrate the merits of a
12 claim that charges for a manufacturer’s
13 product have exceeded the applicable ceil-
14 ing price under this section, and may sub-
15 mit such documents and information to the
16 administrative official or body responsible
17 for adjudicating such claim;

18 “(iv) require that a manufacturer con-
19 duct an audit of a covered entity pursuant
20 to subsection (a)(5)(D) as a prerequisite to
21 initiating administrative dispute resolution
22 proceedings against a covered entity;

23 “(v) permit the official or body des-
24 ignated under clause (i), at the request of
25 a manufacturer or manufacturers, to con-

1 solidate claims brought by more than one
2 manufacturer against the same covered en-
3 tity where, in the judgment of such official
4 or body, consolidation is appropriate and
5 consistent with the goals of fairness and
6 economy of resources; and

7 “(vi) include provisions and proce-
8 dures to permit multiple covered entities to
9 jointly assert claims of overcharges by the
10 same manufacturer for the same drug or
11 drugs in one administrative proceeding,
12 and permit such claims to be asserted on
13 behalf of covered entities by associations or
14 organizations representing the interests of
15 such covered entities and of which the cov-
16 ered entities are members.

17 “(C) FINALITY OF ADMINISTRATIVE RESO-
18 LUTION.—The administrative resolution of a
19 claim or claims under the regulations promul-
20 gated under subparagraph (A) shall be a final
21 agency decision and shall be binding upon the
22 parties involved, unless invalidated by an order
23 of a court of competent jurisdiction.

24 “(4) AUTHORIZATION OF APPROPRIATIONS.—

25 There are authorized to be appropriated to carry out

1 this subsection, such sums as may be necessary for
2 fiscal year 2010, and each succeeding fiscal year.”.

3 (f) CONFORMING AMENDMENTS.—

4 (1) SOCIAL SECURITY ACT.—Section 1927 of
5 the Social Security Act (42 U.S.C. 1396r–8), is
6 amended—

7 (A) in subsection (a)(5)—

8 (i) in subparagraph (A), by striking
9 “covered outpatient drugs” and inserting
10 “covered drugs (as defined in section
11 340B(b)(2) of the Public Health Service
12 Act)”;

13 (ii) by striking subparagraph (D); and

14 (iii) by redesignating subparagraph
15 (E) as subparagraph (D);

16 (B) in subsection (c)(1)(C)(i), by redesignig-
17 nating subclauses (II) through (IV) as sub-
18 clauses (III) through (V), respectively and by
19 inserting after subclause (I) the following new
20 subclause:

21 “(II) any prices charged for a
22 covered drug (as defined in section
23 340B(b)(2) of the Public Health Serv-
24 ice Act);”; and

25 (C) in subsection (k)(1)—

1 (i) in subparagraph (A), by striking
 2 “subparagraph (B)” and inserting “sub-
 3 paragraphs (B) and (D)”; and

4 (ii) by adding at the end the following
 5 new subparagraph:

6 “(D) CALCULATION FOR COVERED
 7 DRUGS.—With respect to a covered drug (as de-
 8 fined in section 340B(b)(2) of the Public
 9 Health Service Act), the average manufacturer
 10 price shall be determined in accordance with
 11 subparagraph (A) except that, in the event a
 12 covered drug is not distributed to the retail
 13 pharmacy class of trade, it shall mean the aver-
 14 age price paid to the manufacturer for the drug
 15 in the United States by wholesalers for drugs
 16 distributed to the acute care class of trade,
 17 after deducting customary prompt pay dis-
 18 counts. The Secretary shall establish a mecha-
 19 nism for collecting the necessary data for the
 20 acute care class of trade from manufacturers.”.

21 (2) PUBLIC HEALTH SERVICE ACT.—Section
 22 340B(a) of such Act (42 U.S.C. 256b(a)) is amend-
 23 ed—

24 (A) in subsection (a)(1), by adding at the
 25 end the following: “Each such agreement shall

1 require that the manufacturer furnish the Sec-
2 retary with reports, on a quarterly basis, of the
3 price for each covered drug subject to the
4 agreement that, according to the manufacturer,
5 represents the maximum price that covered en-
6 tities may permissibly be required to pay for the
7 drug (referred to in this section as the ‘ceiling
8 price’), and shall require that the manufacturer
9 offer each covered entity covered drugs for pur-
10 chase at or below the applicable ceiling price if
11 such drug is made available to any other pur-
12 chaser at any price.”; and

13 (B) in the first sentence of subsection
14 (a)(5)(E), as so redesignated by subsection
15 (c)(2), by inserting “after an audit as described
16 in subparagraph (D), and” after “finds,”.

17 **SEC. 3. EFFECTIVE DATES.**

18 (a) **IN GENERAL.**—The amendments made by this
19 Act shall take effect on January 1, 2010, and shall apply
20 to drugs purchased on or after January 1, 2010.

21 (b) **EFFECTIVENESS.**—The amendments made by
22 this Act shall be effective, and shall be taken into account
23 in determining whether a manufacturer is deemed to meet
24 the requirements of section 340B(a) of the Public Health
25 Service Act (42 U.S.C. 256b(a)) and of section 1927(a)(5)

1 of the Social Security Act (42 U.S.C. 1396r-8(a)(5)), not-
2 withstanding any other provision of law.

○