PAYMENT BASED ON COMPETITION.—

(1) IN GENERAL.—Title XVIII is amended by inserting after section 1847A, as added by subsection (c), the following new section:

‘‘COMPETITIVE ACQUISITION OF OUTPATIENT DRUGS AND BIOLOGICALS
‘‘SEC. 1847B. (a) IMPLEMENTATION OF COMPETITIVE ACQUISITION.—
‘‘(1) IMPLEMENTATION OF PROGRAM.—
‘‘(A) IN GENERAL.—The Secretary shall establish and implement a competitive acquisition program under which—

‘‘(i) competitive acquisition areas are established for contract award purposes for acquisition of and payment for categories of competitively biddable drugs and biologicals (as defined in paragraph (2)) under this part;
‘‘(ii) each physician is given the opportunity annually to elect to obtain drugs and biologicals under the program, rather than under section 1847A; and
‘‘(iii) each physician who elects to obtain drugs and biologicals under the program makes an annual selection under paragraph (5) of the contractor through which drugs and biologicals within a category of drugs and biologicals will be acquired and delivered to the physician under this part.

This section shall not apply in the case of a physician who elects section 1847A to apply.

‘‘(B) IMPLEMENTATION.—For purposes of implementing the program, the Secretary shall establish categories of competitively biddable drugs and biologicals. The Secretary shall phase in the program with respect to those categories beginning in 2006 in such manner as the Secretary determines to be appropriate.

‘‘(C) WAIVER OF CERTAIN PROVISIONS.—In order to promote competition, in carrying out the program the Secretary may waive such provisions of the Federal Acquisition Regulation as are necessary for the efficient implementation of this section, other than provisions relating to confidentiality of information and such other provisions as the Secretary determines appropriate.

‘‘(D) EXCLUSION AUTHORITY.—The Secretary may exclude competitively biddable drugs and biologicals (including a class of such drugs and biologicals) from the competitive bidding system under this section if the application of competitive bidding to such drugs or biologicals—

‘‘(i) is not likely to result in significant savings;
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(ii) is likely to have an adverse impact on access
to such drugs or biologicals.

(2) COMPETITIVELY BIDDABLE DRUGS AND BIOLOGICALS AND
PROGRAM DEFINED.—For purposes of this section—

(A) COMPETITIVELY BIDDABLE DRUGS AND BIOLOGICALS
DEFINED.—The term ‘competitively biddable drugs and
biologicaes’ means a drug or biological described in section
1842(o)(1)(C) and furnished on or after January 1, 2006.

(B) PROGRAM.—The term ‘program’ means the
competitive acquisition program under this section.

(C) COMPETITIVE ACQUISITION AREA; AREA.—The terms
‘competitive acquisition area’ and ‘area’ mean an appropriate
geographic region established by the Secretary under
the program.

(D) CONTRACTOR.—The term ‘contractor’ means an
entity that has entered into a contract with the Secretary
under this section.

(3) APPLICATION OF PROGRAM PAYMENT METHODOLOGY.—

(A) IN GENERAL.—With respect to competitively biddable
drugs and biologicals which are supplied under the
program in an area and which are prescribed by a physician
who has elected this section to apply—

(i) the claim for such drugs and biologicals shall
be submitted by the contractor that supplied the drugs
and biologicals;

(ii) collection of amounts of any deductible and
coinsurance applicable with respect to such drugs and
biologicaes shall be the responsibility of such contractor
and shall not be collected unless the drug or biological
is administered to the individual involved; and

(iii) the payment under this section (and related
amounts of any applicable deductible and coinsurance)
for such drugs and biologicals—

(I) shall be made only to such contractor;

and

(II) shall be conditioned upon the administration
of such drugs and biologicals.

(B) PROCESS FOR ADJUSTMENTS.—The Secretary shall
provide a process for adjustments to payments in the case
in which payment is made for drugs and biologicals which
were billed at the time of dispensing but which were not
actually administered.

(C) INFORMATION FOR PURPOSES OF COST-SHARING.—
The Secretary shall provide a process by which physicians
submit information to contractors for purposes of the collection
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of any applicable deductible or coinsurance amounts under subparagraph (A)(ii).

"(4) CONTRACT REQUIRED.—Payment may not be made under this part for competitively biddable drugs and biologicals prescribed by a physician who has elected this section to apply within a category and a competitive acquisition area with respect to which the program applies unless—

"(A) the drugs or biologicals are supplied by a contractor with a contract under this section for such category of drugs and biologicals and area; and

"(B) the physician has elected such contractor under paragraph (5) for such category and area.

"(5) CONTRACTOR SELECTION PROCESS.—

"(A) ANNUAL SELECTION.—

"(i) IN GENERAL.—The Secretary shall provide a process for the selection of a contractor, on an annual basis and in such exigent circumstances as the Secretary may provide and with respect to each category of competitively biddable drugs and biologicals for an area by selecting physicians.

"(ii) TIMING OF SELECTION.—The selection of a contractor under clause (i) shall be made at the time of the election described in section 1847A(a) for this section to apply and shall be coordinated with agreements entered into under section 1842(h).

"(B) INFORMATION ON CONTRACTORS.—The Secretary shall make available to physicians on an ongoing basis, through a directory posted on the Internet website of the Centers for Medicare & Medicaid Services or otherwise and upon request, a list of the contractors under this section in the different competitive acquisition areas.

"(C) SELECTING PHYSICIAN DEFINED.—For purposes of this section, the term ‘selecting physician’ means, with respect to a contractor and category and competitive acquisition area, a physician who has elected this section to apply and has selected to apply under this section such contractor for such category and area.

"(b) PROGRAM REQUIREMENTS.—

"(1) CONTRACT FOR COMPETITIVELY BIDDABLE DRUGS AND BIOLOGICALS.—The Secretary shall conduct a competition among entities for the acquisition of competitively biddable drugs and biologicals. Notwithstanding any other provision of this title, in the case of a multiple source drug, the Secretary shall conduct such competition among entities for the acquisition of at least one competitively biddable drug and biological within each billing and payment code within each category
for each competitive acquisition area.

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(2) CONDITIONS FOR AWARDING CONTRACT.—

(A) IN GENERAL.—The Secretary may not award a contract to any entity under the competition conducted in a competitive acquisition area pursuant to paragraph (1) with respect to the acquisition of competitively biddable drugs and biologicals within a category unless the Secretary finds that the entity meets all of the following with respect to the contract period involved:

(i) CAPACITY TO SUPPLY COMPETITIVELY BIDDABLE DRUG OR BIOLOGICAL WITHIN CATEGORY.—

(I) IN GENERAL.—The entity has sufficient arrangements to acquire and to deliver competitively biddable drugs and biologicals within such category in the area specified in the contract.

(II) SHIPMENT METHODOLOGY.—The entity has arrangements in effect for the shipment at least 5 days each week of competitively biddable drugs and biologicals under the contract and for the timely delivery (including for emergency situations) of such drugs and biologicals in the area under the contract.

(ii) QUALITY, SERVICE, FINANCIAL PERFORMANCE AND SOLVENCY STANDARDS.—The entity meets quality, service, financial performance, and solvency standards specified by the Secretary, including—

(I) the establishment of procedures for the prompt response and resolution of complaints of physicians and individuals and of inquiries regarding the shipment of competitively biddable drugs and biologicals; and

(II) a grievance and appeals process for the resolution of disputes.

(B) ADDITIONAL CONSIDERATIONS.—The Secretary may refuse to award a contract under this section, and may terminate such a contract, with an entity based upon—

(i) the suspension or revocation, by the Federal Government or a State government, of the entity’s license for the distribution of drugs or biologicals (including controlled substances); or

(ii) the exclusion of the entity under section 1128 from participation under this title.

(C) APPLICATION OF MEDICARE PROVIDER OMBUDSMAN.—

For provision providing for a program-wide Medicare Provider Ombudsman to review complaints, see section 1868(b), as added by section 923 of the Medicare Prescription
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‘‘(3) AWARDING MULTIPLE CONTRACTS FOR A CATEGORY AND AREA.—The Secretary may limit (but not below 2) the number of qualified entities that are awarded such contracts for any category and area. The Secretary shall select among qualified entities based on the following:

‘‘(A) The bid prices for competitively biddable drugs and biologicals within the category and area.

‘‘(B) Bid price for distribution of such drugs and biologicals.

‘‘(C) Ability to ensure product integrity.

‘‘(D) Customer service.

‘‘(E) Past experience in the distribution of drugs and biologicals, including controlled substances.

‘‘(F) Such other factors as the Secretary may specify.

‘‘(4) TERMS OF CONTRACTS.—

‘‘(A) IN GENERAL.—A contract entered into with an entity under the competition conducted pursuant to paragraph (1) is subject to terms and conditions that the Secretary may specify consistent with this section.

‘‘(B) PERIOD OF CONTRACTS.—A contract under this section shall be for a term of 3 years, but may be terminated by the Secretary or the entity with appropriate, advance notice.

‘‘(C) INTEGRITY OF DRUG AND BIOLOGICAL DISTRIBUTION SYSTEM.—A contractor (as defined in subsection (a)(2)(D)) shall—

‘‘(i) acquire all drug and biological products it distributes directly from the manufacturer or from a distributor that has acquired the products directly from the manufacturer; and

‘‘(ii) comply with any product integrity safeguards as may be determined to be appropriate by the Secretary. Nothing in this subparagraph shall be construed to relieve or exempt any contractor from the provisions of the Federal Food, Drug, and Cosmetic Act that relate to the wholesale distribution of prescription drugs or biologicals.

‘‘(D) COMPLIANCE WITH CODE OF CONDUCT AND FRAUD AND ABUSE RULES.—Under the contract—

‘‘(i) the contractor shall comply with a code of conduct, specified or recognized by the Secretary, that includes standards relating to conflicts of interest; and

‘‘(ii) the contractor shall comply with all applicable provisions relating to prevention of fraud and abuse, including compliance with applicable guidelines of the Department of Justice and the Inspector General of
the Department of Health and Human Services.

“(E) DIRECT DELIVERY OF DRUGS AND BIOLOGICALS TO PHYSICIANS.—Under the contract the contractor shall only supply competitively biddable drugs and biologicals directly to the selecting physicians and not directly to individuals, except under circumstances and settings where an individual currently receives a drug or biological in the individual’s home or other non-physician office setting as the Secretary may provide. The contractor shall not deliver drugs and biologicals to a selecting physician except upon receipt of a prescription for such drugs and biologicals, and such necessary data as may be required by the Secretary to carry out this section. This section does not—

“(i) require a physician to submit a prescription for each individual treatment; or

“(ii) change a physician’s flexibility in terms of writing a prescription for drugs or biologicals for a single treatment or a course of treatment.

“(5) PERMITTING ACCESS TO DRUGS AND BIOLOGICALS.—The Secretary shall establish rules under this section under which drugs and biologicals which are acquired through a contractor under this section may be used to resupply inventories of such drugs and biologicals which are administered consistent with safe drug practices and with adequate safeguards against fraud and abuse. The previous sentence shall apply if the physicians can demonstrate to the Secretary all of the following:

“(A) The drugs or biologicals are required immediately.

“(B) The physician could not have reasonably anticipated the immediate requirement for the drugs or biologicals.

“(C) The contractor could not deliver to the physician the drugs or biologicals in a timely manner.

“(D) The drugs or biologicals were administered in an emergency situation.

“(6) CONSTRUCTION.—Nothing in this section shall be construed as waiving applicable State requirements relating to licensing of pharmacies.

“(c) BIDDING PROCESS.—

“(1) IN GENERAL.—In awarding a contract for a category of drugs and biologicals in an area under the program, the Secretary shall consider with respect to each entity seeking to be awarded a contract the bid price and the other factors referred to in subsection (b)(3).

“(2) BID DEFINED.—In this section, the term ‘bid’ means an offer to furnish a competitively biddable drug or biological for a particular price and time period.
“(3) BIDDING ON A NATIONAL OR REGIONAL BASIS.—Nothing in this section shall be construed as precluding a bidder from bidding for contracts in all areas of the United States or as requiring a bidder to submit a bid for all areas of the United States.

“(4) UNIFORMITY OF BIDS WITHIN AREA.—The amount of the bid submitted under a contract offer for any competitively biddable drug or biological for an area shall be the same for that drug or biological for all portions of that area.

“(5) CONFIDENTIALITY OF BIDS.—The provisions of subparagraph (D) of section 1927(b)(3) shall apply to periods during which a bid is submitted with respect to a competitively biddable drug or biological under this section in the same manner as it applies to information disclosed under such section, except that any reference—

‘‘(A) in that subparagraph to a ‘manufacturer or wholesaler’ is deemed a reference to a ‘bidder’ under this section;

‘‘(B) in that section to ‘prices charged for drugs’ is deemed a reference to a ‘bid’ submitted under this section; and

‘‘(C) in clause (i) of that section to ‘this section’, is deemed a reference to ‘part B of title XVIII’.

“(6) INCLUSION OF COSTS.—The bid price submitted in a contract offer for a competitively biddable drug or biological shall—

‘‘(A) include all costs related to the delivery of the drug or biological to the selecting physician (or other point of delivery); and

‘‘(B) include the costs of dispensing (including shipping) of such drug or biological and management fees, but shall not include any costs related to the administration of the drug or biological, or wastage, spillage, or spoilage.

“(7) PRICE ADJUSTMENTS DURING CONTRACT PERIOD; DISCLOSURE OF COSTS.—Each contract awarded shall provide for—

‘‘(A) disclosure to the Secretary the contractor’s reasonable, net acquisition costs for periods specified by the Secretary, not more often than quarterly, of the contract; and

‘‘(B) appropriate price adjustments over the period of the contract to reflect significant increases or decreases in a contractor’s reasonable, net acquisition costs, as so disclosed.

“(d) COMPUTATION OF PAYMENT AMOUNTS.—

‘‘(1) IN GENERAL.—Payment under this section for competitively biddable drugs or biologicals shall be based on bids submitted and accepted under this section for such drugs or biologicals in an area. Based on such bids the Secretary shall
determine a single payment amount for each competitively
biddable drug or biological in the area.

‘‘(2) SPECIAL RULES.—The Secretary shall establish rules
regarding the use under this section of the alternative payment
amount provided under section 1847A to the use of a price
for specific competitively biddable drugs and biologicals in the
following cases:

‘‘(A) NEW DRUGS AND BIOLOGICALS.—A competitively
biddable drug or biological for which a payment and billing
code has not been established.

‘‘(B) OTHER CASES.—Such other exceptional cases as
the Secretary may specify in regulations.

‘‘(e) COST-SHARING.—

‘‘(1) APPLICATION OF COINSURANCE.—Payment under this
section for competitively biddable drugs and biologicals shall
be in an amount equal to 80 percent of the payment basis
described in subsection (d)(1).

‘‘(2) DEDUCTIBLE.—Before applying paragraph (1), the individual
shall be required to meet the deductible described in
section 1833(b).

‘‘(3) COLLECTION.—Such coinsurance and deductible shall
be collected by the contractor that supplies the drug or
biological involved. Subject to subsection (a)(3)(B), such coinsurance
and deductible may be collected in a manner similar
to the manner in which the coinsurance and deductible are
collected for durable medical equipment under this part.

‘‘(f) SPECIAL PAYMENT RULES.—

‘‘(1) USE IN EXCLUSION CASES.—If the Secretary excludes
a drug or biological (or class of drugs or biologicals) under
subsection (a)(1)(D), the Secretary may provide for payment
to be made under this part for such drugs and biologicals
(or class) using the payment methodology under section 1847A.

‘‘(2) APPLICATION OF REQUIREMENT FOR ASSIGNMENT.—For
provision requiring assignment of claims for competitively biddable
drugs and biologicals, see section 1842(o)(3).

‘‘(3) PROTECTION FOR BENEFICIARY IN CASE OF MEDICAL
NECESSITY DENIAL.—For protection of individuals against
liability in the case of medical necessity determinations, see

‘‘(g) JUDICIAL REVIEW.—There shall be no administrative or
judicial review under section 1869, section 1878, or otherwise, of—

‘‘(1) the establishment of payment amounts under subsection
(d)(1);

‘‘(2) the awarding of contracts under this section;

‘‘(3) the establishment of competitive acquisition areas
under subsection (a)(2)(C);
“(4) the phased-in implementation under subsection (a)(1)(B);
“(5) the selection of categories of competitively biddable drugs and biologicals for competitive acquisition under such subsection or the selection of a drug in the case of multiple source drugs; or
“(6) the bidding structure and number of contractors selected under this section.”.

(2) REPORT.—Not later than July 1, 2008, the Secretary shall submit to Congress a report on the program conducted under section 1847B of the Social Security Act, as added by paragraph (1). Such report shall include information on savings, reductions in cost-sharing, access to competitively biddable drugs and biologicals, the range of choices of contractors available to physicians, the satisfaction of physicians and of individuals enrolled under this part, and information comparing prices for drugs and biologicals under such section and section 1847A of such Act, as added by subsection (c).